

STIC Search Report Biotech-Chem Library

STIC Database Tracking Number 1775

TO: Ralph J Gitomer Location: REM-3C18

Art Unit: 1655

Search Notes

Monday, December 05, 2005

Case Serial Number: 10/658609

From: Alex Waclawiw

Location: Biotech-Chem Library

Rem 1A71

Phone: 272-2534

Alexandra.waclawiw@uspto.gov

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Scientific and Technical Information Center

SEARCH REQUEST FORM

Requester's Full Name:	P. GITON	nt N	Examiner # :	69630 D	Date: 11/16/05
Art Unit: 1655	Phone Number		Serial N	umber: / 6/	67 9, 609
		Т	Results Format	Preferred (circle): PAPER DISK
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3 C18 - Co ensure an efficient and qual	ity search, please at	tach a copy of the co	ver sheet, claims, ar	nd abstract or fill o	ut the following:
Title of Invention:				·····	
nventors (please provide fu					
Earliest Priority Date:	-			•	· ·
Search Topic: Please provide a detailed statem elected species or structures, ke Define any terms that may have	cut of the search top wwords, synonyms, ac a special meaning.	oic, and describe as sp cranyms, and registry Give examples or rele	evant citations, auth	ors, etc., if known.	
For Sequence Searches Only appropriate serial number.	Please include all p	pertinent information	(parent, child, divisi	onal, or issued pate	nt numbers) along with the
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STAFF USE ON THE	Contact: T	ype of Search.	Ver	idors and cost wh	ere applicable
Saurcher: Alexandra	Waclawiw	NA Sequence (i	#) 363(STN .	Dialog
Technical Inf	o. Specialist et: 303-4491	AA Sequence (#)	Questel/Orbit	Lexis/Nexis
Searcher Location:		Structure (#)		Westlaw	WWW/Internet
Date Searcher Picked Up:	12-1	Bibliographic		In-house sequen	ce systems
Date Completed:	12-5	Litigation		Commercial	Oligomer Score/Length Encode/Transl
Searcher Prep & Review Time: _	12	Fulltext	<u></u>	Other ((specify)
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FILE 'REGISTRY' ENTERED AT 10:07:48 ON 05 DEC 2005
               E SILVER/CN
             1 SEA ABB=ON PLU=ON SILVER/CN
L1
               E GOLD/CN
L2
             1 SEA ABB=ON PLU=ON GOLD/CN
               E IRON/CN
             1 SEA ABB=ON PLU=ON IRON/CN
L3
               E MERCURY/CN
             1 SEA ABB=ON PLU=ON MERCURY/CN
L4
               E NICKEL/CN
             1 SEA ABB=ON PLU=ON NICKEL/CN
L5
               E COPPER/CN
             1 SEA ABB=ON PLU=ON COPPER/CN
L6
               E PLATINUM/CN
             1 SEA ABB=ON PLU=ON PLATINUM/CN
Ь7
               E PALLADIUM/CN
             1 SEA ABB=ON PLU=ON PALLADIUM/CN
rs
               E COBALT/CN
             1 SEA ABB=ON PLU=ON COBALT/CN
1.9
               E IRIDIUM/CN
             1 SEA ABB=ON PLU=ON IRIDIUM/CN
L10
            10 SEA ABB=ON PLU=ON (L1 OR L2 OR L3 OR L4 OR L5 OR L6 OR L7 OR
L11
               L8 OR L9 OR L10)
               SAVE L11 TEMP METALS/A
     FILE 'CAPLUS' ENTERED AT 10:08:59 ON 05 DEC 2005
       1396141 SEA ABB=ON PLU=ON L11
L12
        555952 SEA ABB=ON PLU=ON L12 (L) (ARG OR ANST OR USES)/RL
L13
           216 SEA ABB=ON PLU=ON
                                  (ENZYM? (S)ALTER?(S) METAL#)/BI
L14
            48 SEA ABB=ON PLU=ON (ENZYM? (L)ALTER?(L) METAL#)/OBI
L15
           217 SEA ABB=ON PLU=ON L14 OR L15
L16
            15 SEA ABB=ON PLU=ON (ENZYM? (L) DEPOSIT? (L) METAL#)/OBI
L17
            46 SEA ABB=ON PLU=ON (ENZYM? (S) DEPOSIT? (S) METAL#)/BI
L18
            47 SEA ABB=ON PLU=ON L17 OR L18
L19
            16 SEA ABB=ON PLU=ON L19 AND L13
L20
            2 SEA ABB=ON PLU=ON L16 AND L13
L21
            17 SEA ABB=ON PLU=ON
                                  (L20 OR L21)
L22
          1312 SEA ABB=ON PLU=ON (HER 2 NEU)/BI
L23
             6 SEA ABB=ON PLU=ON L23 AND L13
L24
 FILE 'REGISTRY' ENTERED AT 10:15:12 ON 05 DEC 2005
L25
             1 SEA ABB=ON PLU=ON PEROXIDASE/CN
               E HYDROGEN PEROXIDE/CN
L26
             1 SEA ABB=ON PLU=ON "HYDROGEN PEROXIDE"/CN
               E HYDROQUINONE/CN
             1 SEA ABB=ON PLU=ON HYDROQUINONE/CN
L27
     FILE 'CAPLUS' ENTERED AT 10:16:23 ON 05 DEC 2005
         58320 SEA ABB=ON PLU=ON L25 OR PEROXIDASE/OBI
L28
         99222 SEA ABB=ON PLU=ON L26 OR HYDROGEN PEROXIDE/OBI
L29
         28855 SEA ABB=ON PLU=ON L27 OR HYDROQUINONE/OBI
L30
           107 SEA ABB=ON PLU=ON L28 AND L29 AND L30
L31
            10 SEA ABB=ON PLU=ON L31 AND L13
L32
        539677 SEA ABB=ON PLU=ON ENZYM?/OBI
L33
             5 SEA ABB=ON PLU=ON L32 AND L33
L34
               D SCAN TI
L35
         96721 SEA ABB=ON PLU=ON PROBE#/OBI
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141986 SEA ABB=ON PLU=ON NUCLEIC ACID/OBI
L36
         28293 SEA ABB=ON PLU=ON L35 (L) L36
L37
               E PROBES/CT
         24165 SEA ABB=ON PLU=ON "PROBES (NUCLEIC ACID)"/CT
         24165 SEA ABB=ON PLU=ON L38 AND L37
L39
            11 SEA ABB=ON PLU=ON L39 AND L23
L40
               D SCA TI
             3 SEA ABB=ON PLU=ON L23 AND L28
L41
             2 SEA ABB=ON PLU=ON L23 AND L29
L42
             0 SEA ABB=ON PLU=ON L23 AND L30
L43
            26 SEA ABB=ON PLU=ON L21 OR L24 OR L34 OR L40 OR L41 OR L42
L44
             5 SEA ABB=ON PLU=ON L23 AND (L28 OR L29 OR L30)
L45
            26 SEA ABB=ON PLU=ON L44 OR L45
L46
       1645688 SEA ABB=ON PLU=ON METAL/OBI OR L13
L47
         11370 SEA ABB=ON PLU=ON L47 (L) (ENZYM?/OBI OR L29)
L48
           230 SEA ABB=ON PLU=ON L48 (L) (BIOSENS?/OBI OR REMEDIAT?/OBI)
L49
             4 SEA ABB=ON PLU=ON L49 AND ANTIBOD?/OBI
L50
             5 SEA ABB=ON PLU=ON L49 AND ANTIGEN#/OBI
L51
            30 SEA ABB=ON PLU=ON L51 OR L50 OR L46
L52
               E HAINSFELD J/AU
               E HAINFELD J/AU
           139 SEA ABB=ON PLU=ON HAINFELD J?/AU
L53
            3 SEA ABB=ON PLU=ON L53 AND L23
L54
             3 SEA ABB=ON PLU=ON L53 AND (L16 OR L19)
L55
            2 SEA ABB=ON PLU=ON L53 AND L39
L56
            9 SEA ABB=ON PLU=ON L53 AND ENZYM?/OBI
L57
            12 SEA ABB=ON PLU=ON (L54 OR L55 OR L56 OR L57)
L58
            10 SEA ABB=ON PLU=ON L58 NOT L52
L59
     FILE 'BIOSIS' ENTERED AT 11:11:13 ON 05 DEC 2005
               E HAINFELD J/AU
           140 SEA ABB=ON PLU=ON ("HAINFELD J"/AU OR "HAINFELD J F"/AU OR
L60
               "HAINFELD JAMES"/AU OR "HAINFELD JAMES F"/AU OR "HAINFELD
               JIM"/AU OR "HAINFELF J"/AU)
          2329 SEA ABB=ON PLU=ON HER 2 NEU OR HER 2NEU
L61
             4 SEA ABB=ON PLU=ON L61 AND L60
L62
         119968 SEA ABB=ON PLU=ON METAL#
L63
         312573 SEA ABB=ON PLU=ON L11 OR GOLD OR SILVER OR IRON OR MERCURY
L64
               OR NICKEL OR COPPER OR PLATINUM OR PALLADIUM OR COBALT OR
               IRIDIUM
       1812729 SEA ABB=ON PLU=ON ENZYM?
L65
          82379 SEA ABB=ON PLU=ON L25 OR PEROXIDASE/OBI
L66
             2 SEA ABB=ON PLU=ON L61 AND (L63 OR L64) AND (L65 OR L66)
L67
          65021 SEA ABB=ON PLU=ON (L63 OR L64) AND (L65 OR L66)
L68
          34450 SEA ABB=ON PLU=ON (L63 OR L64) (L) (L65 OR L66)
L69
             O SEA ABB=ON PLU=ON ENZYM (S) (DEPOSIT?) (S) METAL#
L70
             9 SEA ABB=ON PLU=ON ENZYM? (S) (DEPOSIT?) (S) METAL#
L71
          1072 SEA ABB=ON PLU=ON (ENZYM? (L) (DEPOSIT? OR ALTER?) (L)
L72
               METAL#)
             6 SEA ABB=ON PLU=ON (ENZYM? (L) (DEPOSIT? OR ALTER?) (L)
L73
               METAL#)/IT
         161026 SEA ABB=ON PLU=ON PROBE#
L74
             2 SEA ABB=ON PLU=ON L61 AND L74 AND (L63 OR L64)
L75
             74 SEA ABB=ON PLU=ON L61 AND L74
L76
         116540 SEA ABB=ON PLU=ON PEROXIDASE OR HYDROGEN PEROXIDE OR
L77
               HYDROOUINONE OR L25 OR L26 OR L27
             4 SEA ABB=ON PLU=ON L76 AND L77
L78
            41 SEA ABB=ON PLU=ON L60 AND L64
L79
             9 SEA ABB=ON PLU=ON L79 AND (ENZYM? OR L77)
L80
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		Ralph Gitomer 10/658,609
L81		13 SEA ABB=ON PLU=ON L62 OR L80 D QUE L77
L82		12 SEA ABB=ON PLU=ON L67 OR L73 OR L75 OR L78
L83		12 SEA ABB=ON PLU=ON L81 NOT L82
L84	FILE	'CAPLUS, BIOSIS' ENTERED AT 11:31:03 ON 05 DEC 2005 40 DUP REM L52 L82 (2 DUPLICATES REMOVED) ANSWERS '1-30' FROM FILE CAPLUS ANSWERS '31-40' FROM FILE BIOSIS
L85		21 DUP REM L59 L83 (1 DUPLICATE REMOVED) ANSWERS '1-10' FROM FILE CAPLUS ANSWERS '11-21' FROM FILE BIOSIS

=> fil reg
FILE 'REGISTRY' ENTERED AT 11:31:41 ON 05 DEC 2005
USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT.
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Property values tagged with IC are from the ZIC/VINITI data file provided by InfoChem.

STRUCTURE FILE UPDATES: 4 DEC 2005 HIGHEST RN 869277-23-6 DICTIONARY FILE UPDATES: 4 DEC 2005 HIGHEST RN 869277-23-6

New CAS Information Use Policies, enter HELP USAGETERMS for details.

TSCA INFORMATION NOW CURRENT THROUGH JULY 14, 2005

Please note that search-term pricing does apply when conducting SmartSELECT searches.

Structure search iteration limits have been increased. See HELP SLIMITS for details.

REGISTRY includes numerically searchable data for experimental and predicted properties as well as tags indicating availability of experimental property data in the original document. For information on property searching in REGISTRY, refer to:

http://www.cas.org/ONLINE/UG/regprops.html

```
=> d que 111
             1 SEA FILE=REGISTRY ABB=ON PLU=ON SILVER/CN
L1
             1 SEA FILE=REGISTRY ABB=ON PLU=ON GOLD/CN
L2
             1 SEA FILE=REGISTRY ABB=ON PLU=ON IRON/CN
L3
             1 SEA FILE=REGISTRY ABB=ON PLU=ON MERCURY/CN
L4
             1 SEA FILE=REGISTRY ABB=ON PLU=ON NICKEL/CN
L5
             1 SEA FILE=REGISTRY ABB=ON PLU=ON COPPER/CN
L6
            1 SEA FILE=REGISTRY ABB=ON PLU=ON PLATINUM/CN
L7
            1 SEA FILE=REGISTRY ABB=ON PLU=ON PALLADIUM/CN
LB
            1 SEA FILE=REGISTRY ABB=ON PLU=ON COBALT/CN
L9
             1 SEA FILE=REGISTRY ABB=ON PLU=ON IRIDIUM/CN
L10
            10 SEA FILE=REGISTRY ABB=ON PLU=ON (L1 OR L2 OR L3 OR L4 OR L5
L11
               OR L6 OR L7 OR L8 OR L9 OR L10)
```

=> d que 125;d que 126; d que 127 L25 1 SEA FILE=REGISTRY ABB=ON PLU=ON PEROXIDASE/CN

L26 1 SEA FILE=REGISTRY ABB=ON PLU=ON "HYDROGEN PEROXIDE"/CN

1 SEA FILE=REGISTRY ABB=ON PLU=ON HYDROQUINONE/CN

=> fil caplus biosis FILE 'CAPLUS' ENTERED AT 11:32:36 ON 05 DEC 2005 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2005 AMERICAN CHEMICAL SOCIETY (ACS)

FILE 'BIOSIS' ENTERED AT 11:32:36 ON 05 DEC 2005 Copyright (c) 2005 The Thomson Corporation

```
=> d que 184
L1
             1 SEA FILE=REGISTRY ABB=ON PLU=ON SILVER/CN
             1 SEA FILE=REGISTRY ABB=ON PLU=ON GOLD/CN
L2
             1 SEA FILE=REGISTRY ABB=ON PLU=ON IRON/CN
L3
             1 SEA FILE=REGISTRY ABB=ON PLU=ON MERCURY/CN
T.4
             1 SEA FILE=REGISTRY ABB=ON PLU=ON NICKEL/CN
L5
             1 SEA FILE=REGISTRY ABB=ON PLU=ON COPPER/CN
L6
             1 SEA FILE=REGISTRY ABB=ON PLU=ON PLATINUM/CN
L7
            1 SEA FILE=REGISTRY ABB=ON PLU=ON PALLADIUM/CN
L8
             1 SEA FILE=REGISTRY ABB=ON PLU=ON COBALT/CN
L9
             1 SEA FILE=REGISTRY ABB=ON PLU=ON
L10
                                                IRIDIUM/CN
            10 SEA FILE=REGISTRY ABB=ON PLU=ON (L1 OR L2 OR L3 OR L4 OR L5
L11
               OR L6 OR L7 OR L8 OR L9 OR L10)
       1396141 SEA FILE=CAPLUS ABB=ON PLU=ON L11
L12
        555952 SEA FILE=CAPLUS ABB=ON PLU=ON L12 (L) (ARG OR ANST OR
L13
               USES)/RL
           216 SEA FILE=CAPLUS ABB=ON PLU=ON (ENZYM? (S)ALTER?(S) METAL#)/BI
T.14
            48 SEA FILE=CAPLUS ABB=ON PLU=ON (ENZYM? (L) ALTER?(L) METAL#)/OB
L15
           217 SEA FILE=CAPLUS ABB=ON PLU=ON L14 OR L15
L16
             2 SEA FILE=CAPLUS ABB=ON PLU=ON L16 AND L13
L21
L23
          1312 SEA FILE=CAPLUS ABB=ON PLU=ON
                                              (HER 2 NEU)/BI
            6 SEA FILE=CAPLUS ABB=ON PLU=ON L23 AND L13
L24
L25
             1 SEA FILE=REGISTRY ABB=ON PLU=ON PEROXIDASE/CN
             1 SEA FILE=REGISTRY ABB=ON PLU=ON
                                                "HYDROGEN PEROXIDE"/CN
L26
             1 SEA FILE=REGISTRY ABB=ON PLU=ON HYDROQUINONE/CN
L27
L28
         58320 SEA FILE=CAPLUS ABB=ON PLU=ON L25 OR PEROXIDASE/OBI
         99222 SEA FILE=CAPLUS ABB=ON PLU=ON L26 OR HYDROGEN PEROXIDE/OBI
L29
         28855 SEA FILE=CAPLUS ABB=ON PLU=ON L27 OR HYDROQUINONE/OBI
L30
           107 SEA FILE=CAPLUS ABB=ON PLU=ON L28 AND L29 AND L30
L31
            10 SEA FILE=CAPLUS ABB=ON PLU=ON L31 AND L13
L32
        539677 SEA FILE=CAPLUS ABB=ON PLU=ON ENZYM?/OBI
L33
L34
             5 SEA FILE=CAPLUS ABB=ON PLU=ON L32 AND L33
         96721 SEA FILE=CAPLUS ABB=ON PLU=ON PROBE#/OBI
L35
        141986 SEA FILE=CAPLUS ABB=ON PLU=ON NUCLEIC ACID/OBI
L36
L37
         28293 SEA FILE=CAPLUS ABB=ON PLU=ON L35 (L) L36
                                              "PROBES (NUCLEIC ACID) "/CT
L38
         24165 SEA FILE=CAPLUS ABB=ON PLU=ON
L39
         24165 SEA FILE=CAPLUS ABB=ON PLU=ON L38 AND L37
L40
            11 SEA FILE=CAPLUS ABB=ON PLU=ON
                                             L39 AND L23
L41
             3 SEA FILE=CAPLUS ABB=ON PLU=ON
                                              L23 AND L28
             2 SEA FILE=CAPLUS ABB=ON PLU=ON L23 AND L29
L42
L44
            26 SEA FILE=CAPLUS ABB=ON PLU=ON L21 OR L24 OR L34 OR L40 OR
               L41 OR L42
             5 SEA FILE=CAPLUS ABB=ON PLU=ON L23 AND (L28 OR L29 OR L30)
L45
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1.27

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L46
             26 SEA FILE=CAPLUS ABB=ON
                                       PLU=ON L44 OR L45
       1645688 SEA FILE=CAPLUS ABB=ON PLU=ON
                                               METAL/OBI OR L13
L47
                                       PLU=ON L47 (L) (ENZYM?/OBI OR L29)
L48
         11370 SEA FILE=CAPLUS ABB=ON
                                       PLU=ON
                                               L48 (L) (BIOSENS?/OBI OR
L49
            230 SEA FILE=CAPLUS ABB=ON
                REMEDIAT?/OBI)
              4 SEA FILE=CAPLUS ABB=ON
                                       PLU=ON L49 AND ANTIBOD?/OBI
L50
                                       PLU=ON L49 AND ANTIGEN#/OBI
L51
              5 SEA FILE=CAPLUS ABB=ON
             30 SEA FILE=CAPLUS ABB=ON PLU=ON L51 OR L50 OR L46
L52
           2329 SEA FILE=BIOSIS ABB=ON PLU=ON HER 2 NEU OR HER 2NEU
L61
        119968 SEA FILE=BIOSIS ABB=ON PLU=ON
                                               METAL#
L63
        312573 SEA FILE=BIOSIS ABB=ON PLU=ON L11 OR GOLD OR SILVER OR IRON
L64
                OR MERCURY OR NICKEL OR COPPER OR PLATINUM OR PALLADIUM OR
                COBALT OR IRIDIUM
        1812729 SEA FILE=BIOSIS ABB=ON
                                       PLU=ON
                                               ENZYM?
L65
          82379 SEA FILE=BIOSIS ABB=ON
                                       PLU=ON L25 OR PEROXIDASE/OBI
L66
L67
              2 SEA FILE=BIOSIS ABB=ON
                                       PLU=ON L61 AND (L63 OR L64) AND (L65
                OR L66)
                                       PLU=ON
L73
              6 SEA FILE=BIOSIS ABB=ON
                                               (ENZYM? (L) (DEPOSIT? OR
                ALTER?) (L) METAL#)/IT
        161026 SEA FILE=BIOSIS ABB=ON PLU=ON PROBE#
L74
              2 SEA FILE=BIOSIS ABB=ON PLU=ON L61 AND L74 AND (L63 OR L64)
L75
             74 SEA FILE-BIOSIS ABB-ON PLU-ON L61 AND L74
L76
        116540 SEA FILE=BIOSIS ABB=ON PLU=ON PEROXIDASE OR HYDROGEN
L77
                PEROXIDE OR HYDROQUINONE OR L25 OR L26 OR L27
             4 SEA FILE=BIOSIS ABB=ON PLU=ON L76 AND L77
L78
             12 SEA FILE=BIOSIS ABB=ON PLU=ON L67 OR L73 OR L75 OR L78
L82
             40 DUP REM L52 L82 (2 DUPLICATES REMOVED)
L84
=> d que 185
              1 SEA FILE=REGISTRY ABB=ON PLU=ON SILVER/CN
L1
              1 SEA FILE=REGISTRY ABB=ON PLU=ON GOLD/CN
L2
             1 SEA FILE=REGISTRY ABB=ON PLU=ON IRON/CN
L3
L4
             1 SEA FILE=REGISTRY ABB=ON PLU=ON MERCURY/CN
             1 SEA FILE=REGISTRY ABB=ON PLU=ON NICKEL/CN
L5
             1 SEA FILE=REGISTRY ABB=ON PLU=ON COPPER/CN
L6
             1 SEA FILE=REGISTRY ABB=ON PLU=ON PLATINUM/CN
L7
             1 SEA FILE=REGISTRY ABB=ON PLU=ON PALLADIUM/CN
L8
L9
             1 SEA FILE=REGISTRY ABB=ON PLU=ON COBALT/CN
             1 SEA FILE=REGISTRY ABB=ON PLU=ON IRIDIUM/CN
L10
L11
             10 SEA FILE=REGISTRY ABB=ON PLU=ON (L1 OR L2 OR L3 OR L4 OR L5
                OR L6 OR L7 OR L8 OR L9 OR L10)
       1396141 SEA FILE=CAPLUS ABB=ON PLU=ON L11
L12
                                       PLU=ON L12 (L) (ARG OR ANST OR
        555952 SEA FILE=CAPLUS ABB=ON
L13
                USES)/RL
            216 SEA FILE=CAPLUS ABB=ON
                                       PLU=ON
                                                (ENZYM? (S)ALTER?(S) METAL#)/BI
L14
             48 SEA FILE=CAPLUS ABB=ON
                                       PLU=ON
                                                (ENZYM? (L)ALTER?(L) METAL#)/OB
L15
L16
            217 SEA FILE=CAPLUS ABB=ON
                                       PLU=ON
                                               L14 OR L15
             15 SEA FILE=CAPLUS ABB=ON
                                       PLU=ON
                                               (ENZYM? (L) DEPOSIT? (L)
L17
                METAL#)/OBI
L18
             46 SEA FILE=CAPLUS ABB=ON
                                       PLU=ON
                                               (ENZYM? (S) DEPOSIT? (S)
                METAL#)/BI
             47 SEA FILE=CAPLUS ABB=ON
                                       PLU=ON L17 OR L18
L19
L21
             2 SEA FILE=CAPLUS ABB=ON PLU=ON
                                              L16 AND L13
          1312 SEA FILE=CAPLUS ABB=ON PLU=ON
                                               (HER 2 NEU)/BI
L23
             6 SEA FILE=CAPLUS ABB=ON PLU=ON L23 AND L13
L24
             1 SEA FILE=REGISTRY ABB=ON PLU=ON PEROXIDASE/CN
L25
             1 SEA FILE=REGISTRY ABB=ON PLU=ON "HYDROGEN PEROXIDE"/CN
L26
```

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1 SEA FILE=REGISTRY ABB=ON PLU=ON HYDROQUINONE/CN
L27
        58320 SEA FILE=CAPLUS ABB=ON PLU=ON L25 OR PEROXIDASE/OBI
L28
        99222 SEA FILE=CAPLUS ABB=ON PLU=ON L26 OR HYDROGEN PEROXIDE/OBI
L29
        28855 SEA FILE=CAPLUS ABB=ON PLU=ON L27 OR HYDROQUINONE/OBI
L30
           107 SEA FILE=CAPLUS ABB=ON PLU=ON L28 AND L29 AND L30
L31
            10 SEA FILE=CAPLUS ABB=ON PLU=ON L31 AND L13
L32
        539677 SEA FILE=CAPLUS ABB=ON PLU=ON ENZYM?/OBI
L33
             5 SEA FILE=CAPLUS ABB=ON PLU=ON L32 AND L33
L34
        96721 SEA FILE=CAPLUS ABB=ON PLU=ON PROBE#/OBI
L35
        141986 SEA FILE=CAPLUS ABB=ON PLU=ON NUCLEIC ACID/OBI
L36
        28293 SEA FILE=CAPLUS ABB=ON PLU=ON L35 (L) L36
L37
         24165 SEA FILE=CAPLUS ABB=ON PLU=ON "PROBES (NUCLEIC ACID)"/CT
L38
         24165 SEA FILE=CAPLUS ABB=ON PLU=ON L38 AND L37
L39
            11 SEA FILE=CAPLUS ABB=ON PLU=ON L39 AND L23
L40
            3 SEA FILE=CAPLUS ABB=ON PLU=ON L23 AND L28
L41
            2 SEA FILE=CAPLUS ABB=ON PLU=ON L23 AND L29
L42
            26 SEA FILE=CAPLUS ABB=ON PLU=ON L21 OR L24 OR L34 OR L40 OR
L44
               L41 OR L42
             5 SEA FILE=CAPLUS ABB=ON PLU=ON L23 AND (L28 OR L29 OR L30)
L45
            26 SEA FILE=CAPLUS ABB=ON PLU=ON L44 OR L45
L46
       1645688 SEA FILE=CAPLUS ABB=ON PLU=ON METAL/OBI OR L13
L47
         11370 SEA FILE=CAPLUS ABB=ON PLU=ON L47 (L) (ENZYM?/OBI OR L29)
L48
           230 SEA FILE=CAPLUS ABB=ON PLU=ON L48 (L) (BIOSENS?/OBI OR
L49
               REMEDIAT?/OBI)
             4 SEA FILE=CAPLUS ABB=ON PLU=ON L49 AND ANTIBOD?/OBI
L50
             5 SEA FILE=CAPLUS ABB=ON PLU=ON L49 AND ANTIGEN#/OBI
L51
           30 SEA FILE=CAPLUS ABB=ON PLU=ON L51 OR L50 OR L46
L52
          139 SEA FILE=CAPLUS ABB=ON PLU=ON HAINFELD J?/AU
L53
          3 SEA FILE=CAPLUS ABB=ON PLU=ON L53 AND L23
3 SEA FILE=CAPLUS ABB=ON PLU=ON L53 AND (L16 OR L19)
L54
L55
            2 SEA FILE=CAPLUS ABB=ON PLU=ON L53 AND L39
L56
            9 SEA FILE=CAPLUS ABB=ON PLU=ON L53 AND ENZYM?/OBI
L57
           12 SEA FILE=CAPLUS ABB=ON PLU=ON (L54 OR L55 OR L56 OR L57)
L58
           10 SEA FILE=CAPLUS ABB=ON PLU=ON L58 NOT L52
L59
          140 SEA FILE=BIOSIS ABB=ON PLU=ON ("HAINFELD J"/AU OR "HAINFELD
L60
               J F"/AU OR "HAINFELD JAMES"/AU OR "HAINFELD JAMES F"/AU OR
               "HAINFELD JIM"/AU OR "HAINFELF J"/AU)
          2329 SEA FILE=BIOSIS ABB=ON PLU=ON HER 2 NEU OR HER 2NEU
L61
             4 SEA FILE=BIOSIS ABB=ON PLU=ON L61 AND L60
L62
        119968 SEA FILE=BIOSIS ABB=ON PLU=ON METAL#
L63
        312573 SEA FILE=BIOSIS ABB=ON PLU=ON L11 OR GOLD OR SILVER OR IRON
L64
               OR MERCURY OR NICKEL OR COPPER OR PLATINUM OR PALLADIUM OR
               COBALT OR IRIDIUM
L65
       1812729 SEA FILE=BIOSIS ABB=ON PLU=ON ENZYM?
         82379 SEA FILE=BIOSIS ABB=ON PLU=ON L25 OR PEROXIDASE/OBI
L66
             2 SEA FILE=BIOSIS ABB=ON PLU=ON L61 AND (L63 OR L64) AND (L65
L67
               OR L66)
             6 SEA FILE=BIOSIS ABB=ON PLU=ON (ENZYM? (L) (DEPOSIT? OR
L73
               ALTER?) (L) METAL#)/IT
        161026 SEA FILE=BIOSIS ABB=ON PLU=ON PROBE#
L74
             2 SEA FILE=BIOSIS ABB=ON PLU=ON L61 AND L74 AND (L63 OR L64)
L75
            74 SEA FILE=BIOSIS ABB=ON PLU=ON L61 AND L74
L76
        116540 SEA FILE=BIOSIS ABB=ON PLU=ON PEROXIDASE OR HYDROGEN
L77
               PEROXIDE OR HYDROQUINONE OR L25 OR L26 OR L27
L78
             4 SEA FILE=BIOSIS ABB=ON PLU=ON L76 AND L77
            41 SEA FILE=BIOSIS ABB=ON PLU=ON L60 AND L64
L79
            9 SEA FILE=BIOSIS ABB=ON PLU=ON L79 AND (ENZYM? OR L77)
L80
           13 SEA FILE=BIOSIS ABB=ON PLU=ON L62 OR L80
L81
            12 SEA FILE=BIOSIS ABB=ON PLU=ON L67 OR L73 OR L75 OR L78
L82
           12 SEA FILE=BIOSIS ABB=ON PLU=ON L81 NOT L82
L83
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L85

=> d .ca 184 1-30;d ibib ab it 184 31-40;d ibib 185 1-21

L84 ANSWER 1 OF 40 CAPLUS COPYRIGHT 2005 ACS on STN DUPLICATE 1

ACCESSION NUMBER: 2002:446862 CAPLUS

DOCUMENT NUMBER: 137:363817

Gold-facilitated in situ hybridization: a bright-field TITLE:

autometallographic alternative to fluorescence in situ

hybridization for detection of HER-2

/neu gene amplification

Tubbs, Raymond; Pettay, James; Skacel, Marek; Powell, AUTHOR (S):

Richard; Stoler, Mark; Roche, Patrick; Hainfeld, James

Department of Anatomic and Clinical Pathology, CORPORATE SOURCE:

Cleveland Clinic Foundation, Cleveland, OH, USA

SOURCE: American Journal of Pathology (2002), 160(5),

1589-1595

CODEN: AJPAA4; ISSN: 0002-9440

American Society for Investigative Pathology PUBLISHER:

DOCUMENT TYPE: Journal LANGUAGE: English Entered STN: 14 Jun 2002

Fluorescence in situ hybridization (FISH) represents an excellent method AΒ for profiling gene amplification in situ, but correlation with tissue

morphol. is difficult because of dark-field visualization. Validation of a bright-field assay for assessment of HER-2/

neu gene amplification was investigated. Streptavidin-Nanogold was used to generate bright-field gene copy signals using GoldEnhance gold-based autometallog., catalyzed reported deposition, and a biotin-labeled probe. One hundred cases of invasive breast carcinoma were evaluated for which FISH gene copy results, and mRNA and oncoprotein gene expression, were known. Autometallog. signals were gual. evaluable without the use of oil immersion microscopy. Results correlated well with indirect and direct label FISH. Autometallog. gold-based in situ

hybridization represents a promising bright-field assay for the assessment of HER-2/neu gene amplification.

3-1 (Biochemical Genetics) CC

Section cross-reference(s): 9, 14

IT Gene, animal

RL: ANT (Analyte); ANST (Analytical study)

(ERBB2; gold-facilitated in situ hybridization is a bright-field autometallog. alternative to FISH for detection of HER-

2/neu gene amplification)

IT Recombination, genetic

> (amplification; gold-facilitated in situ hybridization is a bright-field autometallog. alternative to FISH for detection of HER-2/neu gene amplification)

IT Microscopy

(bright field; gold-facilitated in situ hybridization is a bright-field autometallog. alternative to FISH for detection of HER-

2/neu gene amplification)

Mammary gland, neoplasm IT

> (carcinoma; gold-facilitated in situ hybridization is a bright-field autometallog. alternative to FISH for detection of HER-

2/neu gene amplification)

IT Human

> (gold-facilitated in situ hybridization is a bright-field autometallog. alternative to FISH for detection of HER-2/

neu gene amplification)

IT Immunoassay (immunohistochem.; gold-facilitated in situ hybridization is a bright-field autometallog. alternative to FISH for detection of HER-2/neu gene amplification) Nucleic acid hybridization IT (in situ, Autometallog. gold-based; gold-facilitated in situ hybridization is a bright-field autometallog. alternative to FISH for detection of HER-2/neu gene amplification) IT Carcinoma (mammary; gold-facilitated in situ hybridization is a bright-field autometallog. alternative to FISH for detection of HER-2/new gene amplification) 7440-57-5, Gold, biological studies IT RL: ARG (Analytical reagent use); BUU (Biological use, unclassified); ANST (Analytical study); BIOL (Biological study); USES (Uses) (GoldEnhance gold-based autometallog.; gold-facilitated in situ hybridization is a bright-field autometallog. alternative to FISH for detection of HER-2/neu gene amplification) REFERENCE COUNT: 43 THERE ARE 43 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT L84 ANSWER 2 OF 40 CAPLUS COPYRIGHT 2005 ACS on STN DUPLICATE 2 ACCESSION NUMBER: 2000:858199 CAPLUS DOCUMENT NUMBER: 135:163056 Chromogenic in situ hybridization: A practical TITLE: alternative for fluorescence in situ hybridization to detect HER-2/neu oncogene amplification in archival breast cancer samples Tanner, Minna; Gancberg, David; Di Leo, Angelo; AUTHOR (S): Larsimont, Denis; Rouas, Ghizlane; Piccart, Martine J.; Isola, Jorma CORPORATE SOURCE: Laboratory of Cancer Genetics, Technology University and University Hospital of Tampere, Tampere, FIN-33101, Finland American Journal of Pathology (2000), 157(5), SOURCE: 1467-1472 CODEN: AJPAA4; ISSN: 0002-9440 American Society for Investigative Pathology PUBLISHER: DOCUMENT TYPE: Journal English LANGUAGE: Entered STN: 07 Dec 2000 ED AB Determination of HER-2/neu oncogene amplification has become necessary for selection of breast cancer patients for trastuzumab (Herceptin) therapy. Fluorescence in situ hybridization (FISH) is currently regarded as a gold standard method for detecting HER-2/neu amplification, but it is not very practical for routine histopathol. labs. We evaluated a new modification of in situ hybridization, the chromogenic in situ hybridization (CISH), which enables detection of HER-2/neu gene copies with conventional peroxidase reaction. Archival formalin-fixed paraffin-embedded tumor tissue sections were pretreated (by heating in a microwave oven and using enzyme digestion) and hybridized with a digoxigenin-labeled DNA probe. The probe was detected with antidigoxigenin fluorescein, anti-fluorescein peroxidase, and diaminobenzidine. Gene copies visualized by CISH could be easily distinguished with a +40 objective in hematoxylin-stained tissue

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sections. HER-2/neu amplification typically
     appeared as large peroxidase-pos. intranuclear gene copy clusters. CISH
     and FISH (according to Vysis, made from frozen pulverized tumor samples)
     correlated well in a series of 157 breast cancers (kappa coefficient, 0.81).
     The few different classifications were mostly because of low-level
     amplifications by FISH that were neg. by CISH and immunohistochem. with
     monoclonal antibody CB-11. We conclude that CISH, using conventional
     bright-field microscopy in evaluation, is a useful alternative for
determination
     of HER-2/neu amplification in
    paraffin-embedded tumor samples, especially for confirming the immunohistochem.
     staining results.
     3-1 (Biochemical Genetics)
     Section cross-reference(s): 14
     Fusion proteins (chimeric proteins)
TΤ
     RL: ARG (Analytical reagent use); THU (Therapeutic use); ANST (Analytical
     study); BIOL (Biological study); USES (Uses)
        (anti-fluorescein antibody fusion with horseradish peroxidase
        ; for chromogenic in situ hybridization detection of HER-
        2/neu oncogene amplification in breast cancer
        samples)
    Gene, animal
IT
     RL: ANT (Analyte); THU (Therapeutic use); ANST (Analytical study); BIOL
     (Biological study); USES (Uses)
        (c-erbB2; chromogenic in situ hybridization detects HER-
        2/neu oncogene amplification in archival breast
        cancer samples)
TT
    neu (receptor)
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (chromogenic in situ hybridization detects HER-2/
        neu oncogene amplification in archival breast cancer samples)
    Probes (nucleic acid)
IT
     RL: ARG (Analytical reagent use); THU (Therapeutic use); ANST (Analytical
     study); BIOL (Biological study); USES (Uses)
        (digoxigenin-labeled; for chromogenic in situ hybridization detection
        of HER-2/new oncogene amplification in
       breast cancer samples)
    Nucleic acid hybridization
IT
        (in situ, CISH (chromogenic in situ hybridization); chromogenic in situ
        hybridization detects HER-2/neu oncogene
        amplification in archival breast cancer samples)
    Mammary gland
TT
        (neoplasm; chromogenic in situ hybridization detects HER-
        2/neu oncogene amplification in archival breast
        cancer samples)
    Antibodies
IT
    RL: ARG (Analytical reagent use); THU (Therapeutic use); ANST (Analytical
     study); BIOL (Biological study); USES (Uses)
        (to digoxigenin and fluorescein; for chromogenic in situ hybridization
        detection of HER-2/neu oncogene
        amplification in breast cancer samples)
    1672-46-4D, Digoxigenin, DNA conjugates
                                               2321-07-5D, Fluorescein,
IT
    anti-digoxigenin antibody conjugates
                                            66836-18-8, Diaminobenzidine
    RL: ARG (Analytical reagent use); THU (Therapeutic use); ANST (Analytical
     study); BIOL (Biological study); USES (Uses)
        (for chromogenic in situ hybridization detection of HER-
        2/neu oncogene amplification in breast cancer
        samples)
     9003-99-0D, Peroxidase, anti-fluorescein antibody
ΙT
    conjugates
```

RL: ARG (Analytical reagent use); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); USES (Uses)

(horseradish; for chromogenic in situ hybridization detection of HER-2/new oncogene amplification in breast

cancer samples)

REFERENCE COUNT:

THERE ARE 16 CITED REFERENCES AVAILABLE FOR THIS 16 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L84 ANSWER 3 OF 40 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2005:972223 CAPLUS

DOCUMENT NUMBER:

143:278678

TITLE:

Mechanism of Action of (-)-Epigallocatechin-3-Gallate: Auto-oxidation-Dependent Inactivation of Epidermal Growth Factor Receptor and Direct Effects on Growth Inhibition in Human Esophageal Cancer KYSE 150 Cells

AUTHOR (S):

Hou, Zhe; Sang, Shengmin; You, Hui; Lee, Mao-Jung; Hong, Jungil; Chin, Khew-Voon; Yang, Chung S.

CORPORATE SOURCE:

Susan Lehman Cullman Laboratory for Cancer Research, Department of Chemical Biology, Ernest Mario School of Pharmacy, Rutgers, The State University of New Jersey,

Piscataway, NJ, USA

SOURCE:

Cancer Research (2005), 65(17), 8049-8056

CODEN: CNREA8; ISSN: 0008-5472

PUBLISHER:

American Association for Cancer Research

DOCUMENT TYPE: Journal English LANGUAGE:

Entered STN: 07 Sep 2005 ED

(-)-Epigallocatechin-3-gallate (EGCG), the principal polyphenol in green ΑB tea, has been shown to inhibit the growth of many cancer cell lines and to suppress the phosphorylation of epidermal growth factor receptor (EGFR). We observed similar effects of EGCG in esophageal squamous cell carcinoma KYSE 150 cells and epidermoid squamous cell carcinoma A431 cells. Pretreatment of KYSE 150 cells with EGCG (20 μ mol/L) for 0.5 to 24 h in HAM's F12 and RPMI 1640 mixed medium at 37, before the addition of EGF, resulted in a decreased level of phosphorylated EGFR (by 32-85%). Prolonged treatment with EGCG (8 or 24 h) also decreased EGFR protein level (both by 80%). EGCG treatment for 24 h also caused decreased signals of HER-2/neu in esophageal adenocarcinoma OE19 cells. These effects of EGCG were prevented or diminished by the addition of superoxide dismutase (SOD, 5 units/mL), or SOD plus catalase (30 units/mL), to the cell culture medium. A similar phenomenon on inactivation of EGFR was observed in A431 cells as well. Under culture conditions for KYSE 150 cells, EGCG was unstable, with a half-life of .apprx.30 min; EGCG dimers and other oxidative products were formed. The presence of SOD in the culture medium stabilized EGCG and increased its half-life to longer than 24 h and some EGCG epimerized to (+)-gallocatechin-3-gallate. A mechanism of superoxide radical-mediated dimerization of EGCG and H2O2 formation is proposed. The stabilization of EGCG by SOD in the culture medium potentiated the activity of EGCG in inhibiting KYSE 150 cell growth. The results suggest that in cell culture conditions, the auto-oxidation of EGCG leads to EGFR inactivation, but the inhibition of cell growth is due to other mechanisms. It remains to be determined whether the presently observed auto-oxidation of EGCG occurs in vivo. In

future studies of EGCG and other polyphenolic compds. in cell culture, SOD may be added to stabilize EGCG and to avoid possible artifacts.

1-6 (Pharmacology) CC

7722-84-1, Hydrogen peroxide, biological TT 9054-89-1, Superoxide dismutase 11062-77-4, Superoxide RL: BSU (Biological study, unclassified); BIOL (Biological study)

(mechanism of antitumor action of epigallocatechin gallate in human esophageal cancer KYSE 150 cells)

REFERENCE COUNT: 36 THERE ARE 36 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L84 ANSWER 4 OF 40 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2005:207047 CAPLUS

DOCUMENT NUMBER: 142:442529

TITLE: Real-time detection of gene expression in cancer cells

using molecular beacon imaging: new strategies for

cancer research

AUTHOR(S): Peng, Xiang-Hong; Cao, Ze-Hong; Xia, Jin-Tang;

Carlson, Grant W.; Lewis, Melinda M.; Wood, William

C.; Yang, Lily

CORPORATE SOURCE: Department of Surgery, Winship Cancer Institute, Emory

University School of Medicine, Atlanta, GA, 30322, USA

SOURCE: Cancer Research (2005), 65(5), 1909-1917

CODEN: CNREA8; ISSN: 0008-5472

PUBLISHER: American Association for Cancer Research

DOCUMENT TYPE: Journal LANGUAGE: English ED Entered STN: 09 Mar 2005

Development of novel approaches for quant. anal. of gene expression in AΒ intact tumor cells should provide new means for cancer detection and for studying the response of cancer cells to biol. and therapeutic reagents. We developed procedures for detecting the levels of expression of multiple genes in fixed as well as viable cells using mol. beacon imaging technol. We found that simultaneous delivery of mol. beacons targeting survivin and cyclin D1 mRNAs produced strong fluorescence in breast cancer but not in normal breast cells. Importantly, fluorescence intensity correlated well with the level of gene expression in the cells detected by real-time reverse transcription-PCR or Western blot anal. We further show that mol. beacons can detect changes of survivin gene expression in viable cancer cells following epidermal growth factor stimulation, docetaxel treatment, and overexpression of p53 gene. Thus, mol. beacon imaging is a simple and specific method for detecting gene expression in cancer cells. It has great potential for cancer detection and drug development.

CC 3-1 (Biochemical Genetics)

Section cross-reference(s): 1, 14

IT Primers (nucleic acid)

Probes (nucleic acid)

RL: ARG (Analytical reagent use); DGN (Diagnostic use); PRP (Properties); ANST (Analytical study); BIOL (Biological study); USES (Uses)

(real-time detection of gene expression in cancer cells using mol.

beacon imaging as new strategy for cancer research)

IT 851060-61-2

RL: ARG (Analytical reagent use); DGN (Diagnostic use); PRP (Properties); ANST (Analytical study); BIOL (Biological study); USES (Uses)

(nucleotide sequence, HER-2/neu;

real-time detection of gene expression in cancer cells using mol.

beacon imaging as new strategy for cancer research)

REFERENCE COUNT: 24 THERE ARE 24 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L84 ANSWER 5 OF 40 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2005:100736 CAPLUS

DOCUMENT NUMBER: 143:38798

TITLE: Copy number analysis of c-erb-B2 (HER-

2/neu) and topoisomerase II α

genes in breast carcinoma by quantitative real-time

polymerase chain reaction using hybridization probes

and fluorescence in situ hybridization

Murthy, Sabita K.; Magliocco, Anthony M.; Demetrick, AUTHOR (S):

Douglas J.

CORPORATE SOURCE: Department of Pathology, Calgary Laboratory Services,

The University of Calgary, Calgary, AB, Can.

Archives of Pathology & Laboratory Medicine (2005), SOURCE:

129(1), 39-46

CODEN: APLMAS; ISSN: 0003-9985 College of American Pathologists

PUBLISHER:

DOCUMENT . TYPE:

Journal English

LANGUAGE:

4

Entered STN: 07 Feb 2005 ED

The Topoisomerase II α (TOP2A) protein is the target of the AB anthracycline class of chemotherapeutic agents. TOP2A is frequently coamplified with c-erb-B2 and consequently might be a prognostic and/or predictive factor for breast cancer patients when anthracycline-based chemotherapy is a consideration. A total of 20% to 35% of breast carcinomas show amplification of the erb-B2 gene, some of which also have coamplification of the TOP2A gene. Investigation of the prognostic or predictive significance of these gene amplifications requires a reliable and sensitive method for the measurement of gene copy number in clin. tumor samples. To assess 2 different assay methods that might allow accurate, reproducible, quant., and high-throughput estimation of gene copy number in fresh,

frozen, or paraffin-embedded breast cancer specimens. We developed an assay and analyzed the gene copy nos. of the erb-B2 and TOP2A genes in 8 breast cancer cell lines, 6 fresh frozen samples, and 38 paraffin-embedded breast tumor specimens by a novel real-time polymerase chain reaction (PCR) assay using hybridization probes. The results were compared with standard fluorescence in situ hybridization. We discovered a 100% concordance between assessment of gene copy number of erb-B2 and TOP2A, and between quant. PCR and fluorescence in situ hybridization (FISH). Quant. PCR also had the addnl. feature of uncovering an erb-B2 gene polymorphism. Finally, we observed that TOP2A amplification only occurred in conjunction with erb-B2 amplification in our paraffin-embedded cases of invasive breast carcinoma and that this event was present in 5 (42%) of 12 erb-B2 amplified cases. We conclude that the potentially automatic, real-time PCR anal. using hybridization probes is an efficient method to perform copy number anal., with results that appear identical to the FISH technique and with the benefit of identifying HER-2 polymorphisms.

3-1 (Biochemical Genetics) CC

Section cross-reference(s): 13, 14

Probes (nucleic acid)

RL: BSU (Biological study, unclassified); BUU (Biological use, unclassified); BIOL (Biological study); USES (Uses)

(copy number anal. of c-erb-B2 and topoisomerase $\text{II}\alpha$ genes in breast carcinoma by quant. real-time PCR)

REFERENCE COUNT:

THERE ARE 50 CITED REFERENCES AVAILABLE FOR THIS 50 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L84 ANSWER 6 OF 40 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

2004:606388 CAPLUS

DOCUMENT NUMBER:

141:119769

TITLE:

Multi-layered electrochemical microfluidic sensor

comprising reagent on porous layer

INVENTOR (S):

Rossier, Joeel Stephane; Reymond, Frederic; Morier,

Patrick

PATENT ASSIGNEE(S):

Diagnoswiss S.A., Switz. PCT Int. Appl., 52 pp.

SOURCE:

Page 13

12/05/2005 Searched by Alex Waclawiw

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

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APPLICATION NO.
                     KIND DATE
    PATENT NO.
                     ----
                      A1 20040729 WO 2004-EP1013 20040114
    _____
    WO 2004062801
       W: AE, AE, AG, AL, AM, AM, AM, AT, AT, AU, AU, AZ, AZ, BA, BB,
           BG, BG, BR, BR, BW, BY, BY, BZ, BZ, CA, CH, CN, CN, CO, CO, CR,
           CR, CU, CU, CZ, CZ, DE, DE, DK, DK, DM, DZ, EC, EC, EE, EE, EG,
           ES, ES, FI, FI, GB, GD, GE, GE, GH, GH, GH, GM, HR, HR, HU, HU,
           ID, IL, IN, IS, JP, JP, KE, KE, KG, KG, KP, KP, KP, KR, KR, KZ,
           KZ, KZ, LC, LK, LR, LS, LS, LT, LU, LV, MA, MD, MD, MG, MK, MN,
           MW, MX, MX, MZ
                            20051019
                                      EP 2004-701957
                       Α1
    EP 1585596
        R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
           IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK
PRIORITY APPLN. INFO.:
                                        GB 2003-820 A 20030114
                                        WO 2004-EP1013
                                                          W 20040114
```

ED Entered STN: 29 Jul 2004

- AB Microfluidic electrochem. sensor apparatus and a method for conducting anal. tests with said apparatus for multi-reactant assays. The apparatus of this invention is a multi-layer body made of at least three layers, the 1st one being a polymer layer comprising a microstructure with at least one integrated microelectrode and conductive tracks for connection to an external electrochem. unit, the 2nd one being a nonporous material serving to cover said microstructure so as to enable microfluidic manipulations and the 3rd one being a porous layer such as a membrane or a glass frit, said porous layer comprising at least one reagent to be solubilized upon contact with a test solution and reacting with an analyte present in said solution to form a product that is transported along said microstructure so as to enable electrochem. detection of said analyte. The invention notably enables the performance of multi-reactant assays in a reduced number of steps.
- IC ICM B01L003-00

ICS G01N027-40; G01N027-447; G01N033-487

- CC 9-1 (Biochemical Methods)
- IT Antibodies and Immunoglobulins

Antigens

DNA

Enzymes, uses

Ligands

Oligonucleotides

Peptides, uses

Proteins

RNA

Receptors

RL: ARG (Analytical reagent use); DEV (Device component use); ANST (Analytical study); USES (Uses)

(multi-layered electrochem. microfluidic sensor with reagent on porous layer)

IT 9003-99-0, Peroxidase

RL: ANT (Analyte); ANST (Analytical study)

(horseradish; multi-layered electrochem. microfluidic sensor with reagent on porous layer)

IT 123-31-9, Hydroquinone, uses 7722-84-1,

Hydrogen peroxide, uses 27598-85-2, Aminophenol

RL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses)

(multi-layered electrochem. microfluidic sensor with reagent on porous layer)

7440-22-4, Silver, uses 7440-50-8, Copper, uses IT

7440-57-5, Gold, uses 7783-90-6, Silver chloride, uses

9004-34-6, Cellulose, uses

RL: DEV (Device component use); USES (Uses)

(multi-layered electrochem. microfluidic sensor with reagent on porous layer)

L84 ANSWER 7 OF 40 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

2004:870514 CAPLUS

DOCUMENT NUMBER:

142:19426

TITLE:

بغسق

Electrocatalytic H2O2 amperometric detection using

gold nanotube electrode ensembles

AUTHOR(S):

Delvaux, Marc; Walcarius, Alain; Demoustier-Champagne,

Sophie

CORPORATE SOURCE:

Unite de Physique et de Chimie des Hauts Polymeres, Universite Catholique de Louvain, Louvain-la-Neuve,

B-1348, Belg.

SOURCE:

Analytica Chimica Acta (2004), 525(2), 221-230

CODEN: ACACAM; ISSN: 0003-2670

PUBLISHER:

Elsevier B.V.

DOCUMENT TYPE:

Journal

LANGUAGE:

English

Entered STN: 21 Oct 2004

Arrays of nanoscopic gold tubes were prepared by electroless plating of the AB metal within the pores of nanoporous polycarbonate track-etched membranes. A procedure for fabricating an ensemble of enzyme-modified nanoelectrodes has been developed based on the efficient immobilization of horseradish peroxidase (HRP) to the gold nanotubes array using self-assembled monolayers (mercaptoethylamine or mercaptopropionic acid) as anchoring layers. Hydrogen peroxide (H2O2) was determined electrochem. by using gold nanoelectrode ensembles (NEE) functionalized or not in phosphate buffer solution (PB) with or without a mediator (hydroquinone, H2Q). Bare NEE displays a remarkable sensitivity (14 μA mM-1 in H2Q at -0.1 V vs. Aq/AqC1) compared to a classical gold macroelectrode (0.41 μA mM-1). The gold nanoparticles that form the tubular structure act as excellent catalytic surfaces towards the oxidation and the reduction of H2O2. The HRP modified NEE presents a slightly lower sensitivity (9.5 μA mM-1) than bare NEE. However, this system presents an enhanced limit of detection (up to 4+10-6 M) and a higher selectivity towards the detection of H2O2 over a wide range of potentials. The lifetime, fabrication reproducibility and measurement repeatability of the HRP enzyme electrode were evaluated with satisfactory results.

9-7 (Biochemical Methods)

IT Enzyme electrodes

Microelectrodes

Nanoparticles

Nanotubes

Self-assembled monolayers

(electrocatalytic H2O2 amperometric detection using gold nanotube electrode ensembles)

7722-84-1, Hydrogen peroxide, analysis IT

RL: ANT (Analyte); BSU (Biological study, unclassified); ANST (Analytical study); BIOL (Biological study)

(electrocatalytic H2O2 amperometric detection using gold nanotube electrode ensembles)

IΤ 7440-57-5, Gold, uses

RL: DEV (Device component use); USES (Uses)

(electrocatalytic H2O2 amperometric detection using gold nanotube

electrode ensembles)

IT 123-31-9, Hydroquinone, analysis

RL: ARU (Analytical role, unclassified); ANST (Analytical study) (electron mediator; electrocatalytic H2O2 amperometric detection using gold nanotube electrode ensembles)

IT 9003-99-0, Peroxidase

RL: ARG (Analytical reagent use); PEP (Physical, engineering or chemical process); PYP (Physical process); ANST (Analytical study); PROC (Process); USES (Uses)

(horseradish; electrocatalytic H2O2 amperometric detection using gold nanotube electrode ensembles)

REFERENCE COUNT: 52 THERE ARE 52 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L84 ANSWER 8 OF 40 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2004:175492 CAPLUS

DOCUMENT NUMBER: 140:386578

TITLE: Real-time quantitative PCR of microdissected

paraffin-embedded breast carcinoma: An alternative

method for HER-2/neu

analysis

AUTHOR(S): Gjerdrum, Lise Mette; Sorensen, Boe Sandahl; Kjeldsen,

Eigil; Sorensen, Flemming Brandt; Nexo, Ebba;

Hamilton-Dutoit, Stephen

CORPORATE SOURCE: Institutes of Pathology, Aarhus University Hospital,

Aarhus, Den.

SOURCE: Journal of Molecular Diagnostics (2004), 6(1), 42-51

CODEN: JMDIFP; ISSN: 1525-1578

PUBLISHER: Association for Molecular Pathology

DOCUMENT TYPE: Journal LANGUAGE: English ED Entered STN: 04 Mar 2004

We studied the feasibility of using real-time quant. PCR to determine HER-2 DNA AB amplification and mRNA expression in microdissected formalin-fixed, paraffin-embedded breast tumors and compared this with standard immunohistochem. (IHC) and fluorescent in situ hybridization (FISH) methods. Study cases (27 carcinomas and 3 ductal breast carcinoma in situ (DCIS) cases) showed varying Her-2 expression as determined by IHC (HercepTest). In carcinomas, there was a good correlation between HER-2 DNA amplification and strong HER-2 protein expression detected by FISH and IHC, resp. A single DCIS case was amplified in FISH, but not in IHC. Both HER-2 gene amplification and expression could be quantified in microdissected paraffin-embedded tumors using real-time PCR, DNA and RNA being successfully detected in 146 of 150 (97%) and 141 of 150 (94%) samples, resp. PCR anal. for HER-2 DNA amplification using the LightCycler HER2/neu DNA Quantification kit (Roche Mol. Biochems., Mannheim, Germany) correlated fairly well with IHC and FISH. All IHC HER-2 3+ tumors were amplified according to the kit, as was the FISH-amplified DCIS case. DNA-PCR identified five addnl. tumors as being amplified. Interestingly, all these scored 2+ with the HercepTest, but were neg. using FISH. We believe that real-time quant. PCR anal. of HER-2 DNA amplification following microdissection represents a useful supplementary or perhaps even an alternative technique for establishing HER-2 status in paraffin-embedded tumors.

CC 3-1 (Biochemical Genetics)

Section cross-reference(s): 14

IT Probes (nucleic acid)

RL: ARG (Analytical reagent use); DGN (Diagnostic use); PRP (Properties); ANST (Analytical study); BIOL (Biological study); USES (Uses) (5-labeled with FAM and 3'-labeled with TAMRA; establishing gene HER-2

status in paraffin-embedded breast carcinoma using real-time quant. PCR, and comparison of results to those obtained using standard immunohistochem. and FISH hybridization)

REFERENCE COUNT:

THERE ARE 30 CITED REFERENCES AVAILABLE FOR THIS 3.0 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L84 ANSWER 9 OF 40 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

2003:931173 CAPLUS

DOCUMENT NUMBER:

139:391342

TITLE:

4

Anti-estrogen receptor agents for chemotherapy and

prevention

INVENTOR(S):

Hung, Mien-chie; Lau, Yiu-keung; Wen, Yong

PATENT ASSIGNEE(S):

Board of Regents, the University of Texas System, USA PCT Int. Appl., 104 pp.

SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

TAIL	MI INIC	ICI-IFI I	014.														
	PATENT	NO.			KIN		DATE								D.	ATE	
	WO 200	30970	49							WO 2002-US15109					2	0020	514
	W:	ΑE,	AG,	AL,	AM,	ΑT,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	ΒZ,	CA,	CH,	CN,
		CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,
							IN,										
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	RW	: GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	ŪĠ,	ZM,	ZW,	AM,	AZ,	BY,
		KG,	KZ,	MD,	RU,	ТJ,	TM,	AT,	BE,	CH,	CY,	DE,	DK,	ES,	FI,	FR,	GB,
		GR,	ΙE,	IT,	LU,	MC,	NL,	PT,	SE,	TR,	BF,	ВJ,	CF,	CG,	CI,	CM,	GA,
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	Section				ence	(s):	2.	3, 8	. 14	. 15							
IT							_ ,	- • -	,								
	<pre>T Mammary gland, neoplasm (estrogen receptor pos. and HER-2-neu</pre>																

(estrogen receptor pos. and HER-2-no

neg.; anti-estrogen receptor agents for chemotherapy and prevention)

50-07-7, Mitomycin C 50-18-0, Cyclophosphamide 51-21-8, 5-Fluorouracil 57-22-7, Vincristine 59-05-2, Methotrexate 446-72-0, Genistein TT 518-82-1, Emodin 865-21-4, Vinblastine 1605-68-1, Taxane 7440-06-4, Platinum, biological studies 15663-27-1, Cisplatin 23214-92-8, Doxorubicin 33069-62-4, Paclitaxel 41575-94-4, Carboplatin 56420-45-2, Epirubicin 65271-80-9, Mitoxantrone 71486-22-1, Vinorelbine 95058-81-4, Gemcitabine 114977-28-5, Docetaxel 149286-90-8, RG13022 154361-50-9, Capecitabine 180288-69-1, Herceptin RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(anti-estrogen receptor agents for chemotherapy and prevention) THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: 2 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L84 ANSWER 10 OF 40 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2003:913391 CAPLUS

DOCUMENT NUMBER: 139:393102

TITLE: Magneto-controlled method and system for determination

of an analyte in a liquid medium

INVENTOR(S): Willner, Itamar; Katz, Eugenii; Weizmann, Yossi;

Patolsky, Fernando

PATENT ASSIGNEE(S): Yissum Research Development Company of the Hebrew

University of Jerusalem, Israel

SOURCE: PCT Int. Appl., 65 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA.	PATENT NO.				KIND DATE				APPLICATION NO.						DATE			
	2003								1	WO 2	003-	IL36	9		2	0030	506	
WO	2003	0960	14		A3		2004	0304										
	W:	ΑE,	AG,	AL,	ΑM,	ΑT,	ΑU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	ΒZ,	CA,	CH,	CN,	
		CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	
		GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,	LK,	LR,	
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										WO 2	003-1	IL369	9	V	v 20	0030!	506	

ED Entered STN: 21 Nov 2003

AB The present invention concerns a magneto-controlled method and system for the determination of an analyte in a liquid medium. The method and system of the

invention are based on the use of functionalized magnetic particles, e.g. magnetic particles that carry a recognition agent, such that in the presence of the analyte and under appropriate conditions, a chemical reaction occurs yielding a reaction signal. The reaction signal may be an elec. signal, a colorimetric signal, light emission or the formation of a precipitate In accordance with the invention the reaction is significantly enhanced by inducing rapid vibrations or rotations of the magnetic particles on the barrier surface. NADH, DNA, antibodies, and telomerase were determined by various assays. For the telomerase assay, activated amine-functionalized magnetic particles were reacted with a mercaptohexyl-modified nucleic acid that contained a sequence recognized by telomerase. Cell extract (from patients with lung cancer) was mixed with the magnetic particle reagent in the presence of a mixture of nucleotide dNTP that included biotinylated dUTP. Avidin-horseradish peroxidase was allowed to react with any synthesized telomer chains. Bound peroxidase was determined using naphthoguinone-functionalized magnetite particles and luminol. Rotation of the magnetic particles by means of an external rotating magnet amplified the emitted light intensity.

- IC ICM G01N033-53
- CC 9-1 (Biochemical Methods)

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Section cross-reference(s): 3, 7
    Enzymes, analysis
RL: ANT (Analyte); ARG (Analytical reagent use); BSU (Biological study,
IT
    unclassified); CAT (Catalyst use); ANST (Analytical study); BIOL
     (Biological study); USES (Uses)
        (as recognition agent on magnetic particles or as analyte;
        magneto-controlled method and system using magnetic particles carrying
        recognition agents for determination of analytes in liquid media)
    Enzymes, biological studies
IT
    RL: ARG (Analytical reagent use); BSU (Biological study, unclassified);
    CAT (Catalyst use); ANST (Analytical study); BIOL (Biological study); USES
     (Uses)
        (conjugates, with avidin; magneto-controlled method and system using
        magnetic particles carrying recognition agents for determination of
analytes in
        liquid media)
IT
    Avidins
    RL: ARG (Analytical reagent use); BSU (Biological study, unclassified);
    ANST (Analytical study); BIOL (Biological study); USES (Uses)
        (conjugates, with enzymes; magneto-controlled method and
        system using magnetic particles carrying recognition agents for
determination
        of analytes in liquid media)
     7440-06-4, Platinum, analysis 7440-22-4, Silver,
IT
     analysis 7440-57-5, Gold, analysis
    RL: ARU (Analytical role, unclassified); DEV (Device component use);
    ANST (Analytical study); USES (Uses)
        (electrode; magneto-controlled method and system using magnetic
        particles carrying recognition agents for determination of analytes in
liquid
        media)
    243986-04-1
IT
     RL: ARU (Analytical role, unclassified); FMU (Formation, unclassified);
     PEP (Physical, engineering or chemical process); PYP (Physical process);
     ANST (Analytical study); FORM (Formation, nonpreparative); PROC (Process)
        (formation and precipitation of, in DNA anal. using horseradish
        peroxidase; magneto-controlled method and system using magnetic
        particles carrying recognition agents for determination of analytes in
liquid
        media)
IT
     9003-99-0D, Peroxidase, conjugates with avidin
     RL: ARG (Analytical reagent use); BSU (Biological study, unclassified);
     CAT (Catalyst use); ANST (Analytical study); BIOL (Biological study); USES
     (Uses)
        (magneto-controlled method and system using magnetic particles carrying
        recognition agents for determination of analytes in liquid media)
IT
    7722-84-1, Hydrogen peroxide, analysis
    RL: ARU (Analytical role, unclassified); FMU (Formation, unclassified);
    ANST (Analytical study); FORM (Formation, nonpreparative)
        (magneto-controlled method and system using magnetic particles carrying
        recognition agents for determination of analytes in liquid media)
IT
    123-31-9, Hydroquinone, analysis
     RL: ARU (Analytical role, unclassified); FMU (Formation, unclassified);
    RCT (Reactant); ANST (Analytical study); FORM (Formation, nonpreparative);
    RACT (Reactant or reagent)
        (magneto-controlled method and system using magnetic particles carrying
        recognition agents for determination of analytes in liquid media)
L84 ANSWER 11 OF 40 CAPLUS COPYRIGHT 2005 ACS on STN
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2003:512075 CAPLUS

ACCESSION NUMBER:

DOCUMENT NUMBER: 139:63320

TITLE: Anti-estrogen receptor agents for chemotherapy

INVENTOR(S): Hung, Mien-Chie; Lau, Yiu-Keung; Wen, Yong

PATENT ASSIGNEE(S): US

SOURCE: U.S. Pat. Appl. Publ., 49 pp.

CODEN: USXXCO

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE _____ ---------_____ US 2002-142115 US 2003125265 A1 20030703 20020509 PRIORITY APPLN. INFO.: US 2001-289658P P 20010509

ED Entered STN: 04 Jul 2003

AB Methods and compns. regarding the prevention of ER-pos. cancer and the treatment of ER-pos. HER-2/neu-neg. breast cancer are disclosed. Compns. exhibiting both tyrosine kinase inhibitor activity and anti-estrogen receptor activity are useful in the cancer treatment. Emodin, having tyrosine kinase inhibitory activity and anti-estrogen activity, mediated chemopreventive activity of breast tumor development in transgenic mice.

IC ICM A61K031-7048

ICS A61K031-353; A61K031-12

INCL 514027000; 514456000; 514680000

CC 1-6 (Pharmacology)

Section cross-reference(s): 63

50-07-7, Mitomycin C 50-18-0, Cyclophosphamide 51-21-8, 5-Fluorouracil 57-22-7, Vincristine 59-05-2, Methotrexate 289-95-2D, Pyrimidine, Fluoro derivs. 865-21-4, Vinblastine 7440-06-4D, Platinum, compds. 15663-27-1, Cisplatin 23214-92-8, Doxorubicin 33069-62-4, Paclitaxel 41575-94-4, Carboplatin 56420-45-2, Epirubicin 65271-80-9, Mitoxantrone 71486-22-1, Vinorelbine 95058-81-4, Gemcitabine 114977-28-5, Docetaxel 154361-50-9, Capecitabine 180288-69-1, Herceptin

RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(chemotherapy with; anti-estrogen receptor and tyrosine kinase inhibitor agents for cancer chemotherapy)

L84 ANSWER 12 OF 40 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2003:263082 CAPLUS

DOCUMENT NUMBER: 140:55753

TITLE: Immobilization of horseradish peroxidase to

a nano-Au monolayer modified chitosan-entrapped carbon

paste electrode for the detection of hydrogen

peroxide

AUTHOR(S): Lei, Cun-Xi; Hu, Shun-Qin; Shen, Guo-Li; Yu, Ru-Qin CORPORATE SOURCE: College of Chemistry and Chemical Engineering, State

Key Laboratory for Chemo/Biosensing and Chemometrics,

Hunan University, People's Republic of Changsha,

410082, Peop. Rep. China

SOURCE: Talanta (2003), 59(5), 981-988

CODEN: TLNTA2; ISSN: 0039-9140

PUBLISHER: Elsevier Science B.V.

DOCUMENT TYPE: Journal LANGUAGE: English ED Entered STN: 06 Apr 2003

AB A procedure for fabricating an enzyme electrode has been described based

on the effective immobilization of horseradish peroxidase (HRP) to a nano-scaled particulate gold (nano-Au) monolayer modified chitosan-entrapped carbon paste electrode (CCPE). The high affinity of chitosan entrapped in CCPE for nano-Au associated with its amino groups has been utilized to realize the use of nano-Au as an intermediator to retain high bioactivity of the enzyme. Hydrogen peroxide (H2O2) was determined in the presence of hydroquinone as a mediator to transfer electrons between the electrode and HRP. The HRP immobilized on nano-Au displayed excellent electrocatalytical activity to the reduction of H2O2. The effects of exptl. variables such as the operating potential of the working electrode, mediator concentration and pH of measuring solution were investigated for optimum anal. performance by using an amperometric method. The enzyme electrode provided a linear response to hydrogen peroxide over a concentration range of 1.22+10-5-2.43+10-3 mol 1-1 with a sensitivity of 0.013 A 1mol-1 cm-2 and a detection limit of 6.3 μ mol 1-1 based on signal per noise =3. The apparent Michaelis-Menten constant (Kmapp) for the sensor was found to be 0.36 mmol 1-1. The lifetime, fabrication reproducibility and measurement repeatability were evaluated with satisfactory results. The anal. results of real sample by this sensor were in satisfactory agreement with those of the potassium permanganate titration method. 9-1 (Biochemical Methods) horseradish peroxidase electrode hydrogen peroxide carbon paste chitosan gold Paste electrodes (carbon; immobilization of horseradish peroxidase to a nano-Au monolayer modified chitosan-entrapped carbon paste electrode for detection of hydrogen peroxide) Amperometry Cyclic voltammetry Enzyme electrodes Immobilization, molecular or cellular Michaelis constant рΗ (immobilization of horseradish peroxidase to a nano-Au monolayer modified chitosan-entrapped carbon paste electrode for detection of hydrogen peroxide) Enzymes, uses RL: ARG (Analytical reagent use); DEV (Device component use); PEP (Physical, engineering or chemical process); PYP (Physical process); ANST (Analytical study); PROC (Process); USES (Uses) (immobilized; immobilization of horseradish peroxidase to a nano-Au monolayer modified chitosan-entrapped carbon paste electrode for detection of hydrogen peroxide) Stability (storage, of HRP electrode; immobilization of horseradish peroxidase to a nano-Au monolayer modified chitosan-entrapped carbon paste electrode for detection of hydrogen peroxide) **7440-57-5**, Gold, analysis RL: ARU (Analytical role, unclassified); DEV (Device component use); ANST (Analytical study); USES (Uses) (colloidal; immobilization of horseradish peroxidase to a nano-Au monolayer modified chitosan-entrapped carbon paste electrode for detection of hydrogen peroxide) 9003-99-0, Peroxidase RL: ARG (Analytical reagent use); DEV (Device component use); PEP (Physical, engineering or chemical process); PYP (Physical process); ANST

(horseradish; immobilization of horseradish peroxidase to a

(Analytical study); PROC (Process); USES (Uses)

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nano-Au monolayer modified chitosan-entrapped carbon paste electrode for detection of hydrogen peroxide) 7722-84-1, Hydrogen peroxide, analysis IT RL: ANT (Analyte); ANST (Analytical study) (immobilization of horseradish peroxidase to a nano-Au monolayer modified chitosan-entrapped carbon paste electrode for detection of hydrogen peroxide) 123-31-9, Hydroquinone, analysis TT RL: ARU (Analytical role, unclassified); ANST (Analytical study)

(immobilization of horseradish peroxidase to a nano-Au monolayer modified chitosan-entrapped carbon paste electrode for detection of hydrogen peroxide)

9012-76-4, Chitosan IT

RL: ARU (Analytical role, unclassified); DEV (Device component use); ANST (Analytical study); USES (Uses)

(immobilization of horseradish peroxidase to a nano-Au monolayer modified chitosan-entrapped carbon paste electrode for detection of hydrogen peroxide)

7440-44-0, Carbon, uses IT

> RL: DEV (Device component use); USES (Uses) (paste electrodes; immobilization of horseradish peroxidase to a nano-Au monolayer modified chitosan-entrapped carbon paste

electrode for detection of hydrogen peroxide)

REFERENCE COUNT: 30 THERE ARE 30 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L84 ANSWER 13 OF 40 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

2004:101837 CAPLUS

DOCUMENT NUMBER:

140:298191

TITLE:

Automatic quantification of gene amplification in

clinical samples by IQ-FISH

AUTHOR(S): CORPORATE SOURCE:

SOURCE:

Narath, R.; Loerch, T.; Rudas, M.; Ambros, P. F. CCRI, Children's Cancer Research Institute, St. Anna Kinderspital, Vienna, A-1090, Austria

Cytometry, Part B: Clinical Cytometry (2003), Volume

Date 2004, 57B(1), 15-22

CODEN: CPBCB5

PUBLISHER:

Wiley-Liss, Inc.

DOCUMENT TYPE:

Journal

LANGUAGE:

English

Entered STN: 09 Feb 2004

The reliable detection and quantification of gene amplifications is AΒ crucial to clin. practice. Although there are different detection techniques, the fluorescence in situ hybridization (FISH) method has become highly accepted over past years because it is a reliable, robust, and quick method. Unfortunately, automatic quantification of gene amplification based on fluorescence intensities has not been possible thus far. Because current spot counting methods are reliable only when analyzing low amplification rates, we attempted to establish another method, i.e., to quantify the intensity of different FISH signals using an automatic fluorescence microscopical device on interphase nuclei: interphase quant. FISH (IQ-FISH). We quantified the fluorescence intensities of the differently labeled FISH probes (MYCN and D2Z) hybridized to three different neuroblastoma cell lines, six peripheral blood (PB) samples, 10 spiked PB samples, and nine neuroblastoma samples using the Metafer4 system (MetaSystems, Altlussheim, Germany). To obtain the MYCN copy number per cell, the ratio between the fluorescence intensities of the MYCN gene and reference sequence (D2Z) was calculated For automatic

of the HER-2/neu status in tumor cells,

anal.

labeled FISH probes specific for HER-2/neu and a chromosome 17-specific probe were hybridized to peripheral blood and tumor specimens and analyzed using the automatic device. When measuring the fluorescence intensity per cell for both probe pairs (MYCN/D2Z and HER-2/17p), amplified and non-amplified cells, showed distinct peaks with only little overlap. Whereas normal cells showed a fluorescence ratio peak for MYCN/D2Z between 200 and 800, cells with MYCN amplification clearly exceeded this ratio value (1000 to 25,000). When mixing a varying number of MYCN amplified cells (range 9-91%) to normal PB, the spiked tumor cells could be identified. Even one neuroblastoma tumor cell in 1000 mononucleated cells could reliably be detected using our device. neuroblastoma patient samples, non-amplified cells were distinguished from amplified cells. Automatically and manually counted signals gave matching results in amplified and non-amplified samples. HER-2 /neu-amplified cells were automatically detected in the breast cancer samples analyzed. The automatic measurement of fluorescence signal intensities not only allows a reliable discrimination between non-amplified and amplified cells but also exact quantification of amplified sequences. This is the prerequisite for the following applications: detection of amplified cells in the bone marrow and second-look specimens; comparison between primary and relapse or pre- and post-chemotherapeutic specimens; detection of tumors with focal gene amplification; and quantification of elimination of amplified gene sequences.

CC 3-1 (Biochemical Genetics)
 Section cross-reference(s): 14

IT Probes (nucleic acid)

RL: ARG (Analytical reagent use); DGN (Diagnostic use); ANST (Analytical study); BIOL (Biological study); USES (Uses)

(automatic quantification of gene amplification in clin. samples by IO-FISH)

REFERENCE COUNT:

THERE ARE 13 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L84 ANSWER 14 OF 40 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2002:123042 CAPLUS

DOCUMENT NUMBER: 136:180121

TITLE: Pseudo-metalloproteins, their preparation and use in

biosensors

INVENTOR(S): Lombardi, Angelina; Pavone, Vincenzo

PATENT ASSIGNEE(S): Universita' Degli Studi di Napoli "Federico II", Italy

SOURCE: PCT Int. Appl., 22 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DA					DATE			APPL	ICAT	ION	NO.		D	ATE		
WO 2002	0122	78		A2		2002	0214	,	WO 2	001-	IB14	27		2	0010	809
WO 2002	0122	78		A3		2002	0613									
W:	ΑE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	ΒZ,	CA,	CH,	CN,
	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,
	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	ΚP,	KR,	ΚZ,	LC,	LK,	LR,
	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	ΝŻ,	PL,	PT,
	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	TJ,	TM,	TR,	TT,	TZ,	UA,	ŪĠ,	US,
	UΖ,	VN,	YU,	ZA,	ZW,	AM,	ΑZ,	BY,	KG,	ΚZ,	MD,	RU,	ТJ,	TM		
RW:	GH,	GM,	KΕ,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZW,	AT,	BE,	CH,	CY,
	DE,	DK,	ES,	FI,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE,	TR,	BF,

BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG IT 1317895 В1 20030715 IT 2000-RM454 20000810 CA 2419019 AA 20020214 CA 2001-2419019 20010809 AU 2001076606 **A5** 20020218 AU 2001-76606 20010809 EP 1309612 A2 20030514 EP 2001-954265 20010809 AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR NZ 524643 20040924 NZ 2001-524643 20010809 Α US 2005090649 20050428 US 2003-344329 20010809 **A1** PRIORITY APPLN. INFO.: IT 2000-RM454 A 20000810 WO 2001-IB1427 W 20010809

OTHER SOURCE(S): MARPAT 136:180121

ED Entered STN: 15 Feb 2002

GT

Described herein are Pseudo-metalloproteins (M = metal selected among Fe, AB Mn, Ti, Mo, Co, Ni, Cu, Pd, Pt, Au, Ru, Cr, V, Tb, Yb, Rh, Ir, Os; X1 = antigen, or else a functional group that enables association to a biomol.; X2 = functional group that enables association to an electrode; S1 and S2 = spacer groups made up of a chain of 3-12 atoms of C, N, O, S and corresponding mixts.; all the other substituents have an amino acid nature), their preparation, and electrochem. biosensors containing them. biosensors can be used in various assays such as diagnostic assays, immunodiagnostic assays, determination of pollutants in water, etc. peptide-metal complex, containing Fe3+ as M; substance P sequence as X1; Cys as X2 and C1-4; Gly-Gly as S1 and S2, was prepared The peptides were synthesized on an automatic peptide synthesizer and then complexed with Fe (SO4) 2 (NH4) 2.

ICM C07K014-00 IC

9-1 (Biochemical Methods) CC

Section cross-reference(s): 15, 29, 34, 61

ITAntigens

> Enzyme inhibitors Oligonucleotides

Peptide nucleic acids

Thiols, preparation

RL: ARG (Analytical reagent use); DEV (Device component use); RCT (Reactant); SPN (Synthetic preparation); ANST (Analytical study); PREP (Preparation); RACT (Reactant or reagent); USES (Uses)

(conjugates with peptide-based metal complexes;

pseudo-metalloproteins, preparation and use in biosensors)

Antibodies and Immunoglobulins IT

Antigens

Enzymes, analysis Nucleic acids Peptides, analysis Proteins

RL: ANT (Analyte); BSU (Biological study, unclassified); ANST (Analytical

study); BIOL (Biological study)

(pseudo-metalloproteins, preparation and use in biosensors)

L84 ANSWER 15 OF 40 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2002:755092 CAPLUS

DOCUMENT NUMBER: 137:259635

TITLE: Enzymatic deposition and alteration

of metals

INVENTOR(S): Hainfeld, James F. PATENT ASSIGNEE(S): Nanoprobes, USA

SOURCE: U.S. Pat. Appl. Publ., 18 pp.

CODEN: USXXCO

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 2002142411	A1	20021003	US 2001-822131	20010330
US 6670113	B2	20031230		
PRIORITY APPLN. INFO.:			US 2001-822131	20010330

ED Entered STN: 04 Oct 2002

Disclosed are methods and materials for utilizing enzymes to act on metal ions in solution so that the ions are reduced to metal. Addnl., disclosed is how to use enzymes to accumulate metal particles. The alteration of metal particles by enzymes interacting with the organic shell of the particles is also described. These methods enable a wide range of applications including sensitive detection of genes and proteins, use as probes for microscopy, nanofabrication, biosensors, and remediation.

IC ICM C12P003-00

INCL 435168000

CC 9-2 (Biochemical Methods)

ST enzyme metal deposition biosensor

bioremediation microscopy nanofabrication

IT Transforming proteins

RL: ANT (Analyte); ANST (Analytical study)

(Her 2-neu; enzymic deposition

and alteration of metals in detection of)

IT Remediation

(bioremediation; enzymic deposition and alteration of metals)

IT Biosensors

Microscopy

(enzymic deposition and alteration of

IT Antigens

RL: ANT (Analyte); ANST (Analytical study)
 (enzymic deposition and alteration of
 metals)

IT Eubacteria

Mammary gland, neoplasm

(enzymic deposition and alteration of metals in detection of)

```
IT
    Estrogen receptors
    Progesterone receptors
    RL: ANT (Analyte); ANST (Analytical study)
        (enzymic deposition and alteration of
       metals in detection of)
IT
    7439-89-6, Iron, uses 7439-97-6, Mercury, uses
    7440-02-0, Nickel, uses 7440-22-4, Silver, uses
    7440-50-8, Copper, uses 7440-57-5, Gold, uses
    RL: ARG (Analytical reagent use); BCP (Biochemical process);
    ANST (Analytical study); BIOL (Biological study); PROC (Process);
    USES (Uses)
        (enzymic deposition and alteration of
       metals)
    9001-05-2, Catalase 9003-99-0, Peroxidase
TT
    RL: ARG (Analytical reagent use); BSU (Biological study, unclassified);
    ANST (Analytical study); BIOL (Biological study); USES (Uses)
        (enzymic deposition and alteration of
       metals)
L84 ANSWER 16 OF 40 CAPLUS COPYRIGHT 2005 ACS on STN
ACCESSION NUMBER:
                        2002:941798 CAPLUS
DOCUMENT NUMBER:
                        138:12469
                        Test strip and biosensor incorporating with nanometer
TITLE:
                        metal particles
                        Shen, Thomas Y. S.; Chen, Wen-Chang; Lin, Hong-Ming;
INVENTOR(S):
                        Chuang, Jen-Hung
PATENT ASSIGNEE(S):
                        Apex Biotechnology Corporation, Taiwan
SOURCE:
                        U.S., 13 pp.
                        CODEN: USXXAM
DOCUMENT TYPE:
                        Patent
LANGUAGE:
                        English
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
                                         APPLICATION NO.
    PATENT NO.
                       KIND DATE
                       ____
                                          -----
     _____
                                                                 -----
    US 6491803
                               20021210 US 2001-859371
                                                               20010518
PRIORITY APPLN. INFO.:
                                         US 2001-859371
    Entered STN: 12 Dec 2002
AB
    The present invention relates to a test strip and a biosensor having an
    increased conductivity and a slurry comprising a fiber, metal particles having
    size in nanometer and a bioactive substance. The invention is
    characterized by incorporating metal particles having size in nanometer
    into the reaction layer of test strip and biosensor to increase the
conductivity
    between the reaction layer and the electrodes so that the redox reaction
    can be readily completed and the measurement time can thus be shortened.
    ICM G01N027-327
TC.
    ICS B05D003-00
INCL 204403110; 204403060; 204403100; 204403010; 427002130
    9-1 (Biochemical Methods)
IT
    Enzymes, uses
    RL: ARG (Analytical reagent use); DEV (Device component use); ANST
     (Analytical study); USES (Uses)
        (immobilized; test strip and biosensor incorporating with
       nanometer metal particles)
    Antibodies and Immunoglobulins
IT
      Antigens
      Enzymes, uses
```

RL: ARG (Analytical reagent use); DEV (Device component use); ANST (Analytical study); USES (Uses)

(test strip and biosensor incorporating with nanometer

metal particles)

REFERENCE COUNT:

THERE ARE 8 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L84 ANSWER 17 OF 40 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

2002:213734 CAPLUS

DOCUMENT NUMBER:

136:230708

TITLE:

4.1

Test kits for determining breast cancer prognosis by

HER-2/neu gene

amplification using fluorescent in-situ hybridization

and control cell lines

INVENTOR(S):

Jaffee, Deborah R.; Flom, Kerry J.

PATENT ASSIGNEE(S):

Ventana Medical Systems, Inc., USA

SOURCE:

U.S., 16 pp., Cont.-in-part of U.S. Provisional Ser.

No. 72,574. CODEN: USXXAM

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 6358682	B1	20020319	US 1999-237115	19990126
US 2003059790	A1	20030327	US 2002-77272	20020215
PRIORITY APPLN. INFO.:			US 1998-72574P	P 19980126
			US 1999-237115	A3 19990126

ED Entered STN: 21 Mar 2002

This invention relates to a method, kit and controls for detecting HER-2/neu gene amplification as a predictor of breast cancer recurrence and patient survival. These patients include those who have had primary, invasive localized breast cancer and who are lymph node-neg. The method is a fluorescent in-situ hybridization (FISH) assay using a labeled DNA probe. More specifically, the method involves counting the number of HER-2/neu genes in a tumor cell. A specific embodiment of the invention is a kit that contains a DNA probe and detection reagents that yields a green fluorescent signal at the site of each HER-2/neu gene on a blue fluorescent background of stained nuclear DNA. The kit is provided for

use with 4 µm sections of formalin-fixed, paraffin-embedded human breast cancer tissue on slides. Furthermore, the simultaneous use of another kit containing control cell lines is recommended. The preferred cells lines used for controls are one with high amplification of the HER -2/neu gene, one with non-amplification and one with

low amplification of this gene. Control tumor cell lines with predefined amts. of HER-2/neu gene amplification

include a non-amplified control cell line with a mean of 3 or fewer HER-2/neu genes per cell (wherein all of the

cells are distributed throughout the slide in 3 dimensions), an amplified control cell line with 10 or more such genes/cell and a lower amplified control cell line with 3-10 HER-2/neu genes

per cell. More specifically, these cell lines are ATCC HTB 30 (SK-BR-3), ATCC HTB 132 (MDA-MB468) and ATCC HTB 133 (T-47D). An average of about 10 or more HER-2/neu genes indicates a high

likelihood of cancer recurrence, while an average of about 3 or fewer indicates a low likelihood of cancer recurrence. Typically, 20-40 tumor cells are counted. By determining the genetic nature of the cancer cells,

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appropriate treatment may be utilized.
IC
     ICM C12Q001-68
     ICS C12P019-34
INCL 435006000
     14-1 (Mammalian Pathological Biochemistry)
CC
     Section cross-reference(s): 3
    Animal cell line
TT
        (ATCC HTB 132 (MDA-MB468), as internal standard for HER-2
        /neu gene amplification; test kits for determining breast cancer
        prognosis by HER-2/neu gene amplification
        using fluorescent in-situ hybridization and control cell lines)
    Gene, animal
TТ
    RL: BSU (Biological study, unclassified); PRP (Properties); THU
     (Therapeutic use); BIOL (Biological study); USES (Uses)
        (ERBB2, of human; test kits for determining breast cancer prognosis by
        HER-2/neu gene amplification using
        fluorescent in-situ hybridization and control cell lines)
TT
    Probes (nucleic acid)
     RL: ARG (Analytical reagent use); DGN (Diagnostic use); ANST (Analytical
     study); BIOL (Biological study); USES (Uses)
        (HER-2/new gene amplification using; test
        kits for determining breast cancer prognosis by HER-2/
        neu gene amplification using fluorescent in-situ hybridization
        and control cell lines)
IT
     Prognosis
        (HER-2/neu gene copy number in breast
        cancer; test kits for determining breast cancer prognosis by HER-
        2/neu gene amplification using fluorescent in-situ
        hybridization and control cell lines)
IT
    Human
        (HER-2/neu gene of; test kits for determining
        breast cancer prognosis by HER-2/neu gene
        amplification using fluorescent in-situ hybridization and control cell
        lines)
IT
    Animal cell line
        (SK-BR-3, as internal standard for HER-2/neu
        gene amplification; test kits for determining breast cancer prognosis by
        HER-2/neu gene amplification using
        fluorescent in-situ hybridization and control cell lines)
IT
     Animal cell line
        (T47D, as internal standard for HER-2/neu
        gene amplification; test kits for determining breast cancer prognosis by
        HER-2/neu gene amplification using
        fluorescent in-situ hybridization and control cell lines)
IT
     Test kits
        (breast cancer prognosis using; test kits for determining breast cancer
        prognosis by HER-2/neu gene amplification
        using fluorescent in-situ hybridization and control cell lines)
IT
     Tumor markers
        (breast cancer, HER-2/neu gene as; test
        kits for determining breast cancer prognosis by HER-2/
        neu gene amplification using fluorescent in-situ hybridization
        and control cell lines)
IT
    Diagnosis
        (cancer, breast, HER-2/neu gene as
        prognostic marker in; test kits for determining breast cancer prognosis by
        HER-2/neu gene amplification using
        fluorescent in-situ hybridization and control cell lines)
IT
     Mammary gland, neoplasm
        (determining prognosis of patients with; test kits for determining breast
cancer
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prognosis by HER-2/new gene amplification
        using fluorescent in-situ hybridization and control cell lines)
IT
     Chromosome
        (human 17, 17q11.2q12, HER-2/neu gene
        mapping to; test kits for determining breast cancer prognosis by HER
        -2/neu gene amplification using fluorescent in-situ
        hybridization and control cell lines)
IT
    Nucleic acid hybridization
        (in situ, fluorescence, HER-2/neu gene
        amplification using; test kits for determining breast cancer prognosis by
        HER-2/neu gene amplification using
        fluorescent in-situ hybridization and control cell lines)
IT
     Gene dosage
        (neu gene, determination of, in diagnosis and prognosis of breast cancer;
test
        kits for determining breast cancer prognosis by HER-2/
        neu gene amplification using fluorescent in-situ hybridization
        and control cell lines)
IT
     Statistical analysis
        (of HER-2/new gene amplification in
        breast tissue samples, for breast cancer prognosis determination; test kits
for
        determining breast cancer prognosis by HER-2/neu
        gene amplification using fluorescent in-situ hybridization and control
        cell lines)
     Genetic mapping
IT
        (of HER-2/neu gene, to human chromosome
        17; test kits for determining breast cancer prognosis by HER-
        2/neu gene amplification using fluorescent in-situ
        hybridization and control cell lines)
IT
    Microscopes
        (slides, breast cancer tissues samples fixed on; test kits for determining
        breast cancer prognosis by HER-2/neu gene
        amplification using fluorescent in-situ hybridization and control cell
        lines)
IT
     403638-47-1
                   403638-48-2
     RL: PRP (Properties)
        (unclaimed nucleotide sequence; test kits for determining breast cancer
        prognosis by HER-2/neu gene amplification
        using fluorescent in-situ hybridization and control cell lines)
REFERENCE COUNT:
                               THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS
                               RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
L84 ANSWER 18 OF 40 CAPLUS COPYRIGHT 2005 ACS on STN
                         2002:604867 CAPLUS
ACCESSION NUMBER:
DOCUMENT NUMBER:
                         138:50447
TITLE:
                         Evaluation of HER-2/neu
                         gene amplification and overexpression: comparison of
                         frequently used assay methods in a molecularly
                         characterized cohort of breast cancer specimens
                         Press, Michael F.; Slamon, Dennis J.; Flom, Kerry J.;
AUTHOR (S):
                         Park, Jinha; Zhou, Jian-Yuan; Bernstein, Leslie
                         Breast Cancer Research Program of the Lee Breast
CORPORATE SOURCE:
                         Center, Department of Pathology and Department of
                         Preventive Medicine, Norris Comprehensive Cancer
                         Center, University of Southern California, Los
                         Angeles, CA, USA
SOURCE:
                         Journal of Clinical Oncology (2002), 20(14), 3095-3105
                         CODEN: JCONDN; ISSN: 0732-183X
                         Lippincott Williams & Wilkins
PUBLISHER:
```

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Journal
DOCUMENT TYPE:
                         English
LANGUAGE:
     Entered STN: 14 Aug 2002
ED
     One hundred seventeen breast cancer specimens with known HER-
AB
     2/neu amplification and overexpression status were
     assayed with four different immunohistochem. assays and two different
     fluorescence in situ hybridization (FISH) assays. The accuracy of the
     FISH assays for HER-2/neu gene amplification
     was high, 97.4% for the Vysis PathVision assay and 95.7% for the the
     Ventana INFORM assay. The immunohistochem. assay with the highest
     accuracy for HER-2/neu overexpression was
     obtained with R60 polyclonal antibody (96.6%), followed by
     immunohistochem. assays performed with 10H8 monoclonal antibody (95.7%),
     the Ventana CB11 monoclonal antibody (89.7%), and the DAKO HercepTest
     (88.9%). Only the sensitivities, and therefore, overall accuracy, of the
     DAKO Herceptest and Ventana CB11 immunohistochem. assays were
     significantly different from the more sensitive FISH assay.
     these findings, the FISH assays were highly accurate, with
     immunohistochem. assays performed with R60 and 10H8 nearly as accurate.
     The DAKO HercepTest and the Ventana CB11 immunohistochem. assay were
     statistically significantly different from the Vysis FISH assay in
     evaluating these previously characterized breast cancer specimens.
     3-1 (Biochemical Genetics)
CC
     Section cross-reference(s): 6, 13, 15
     Centromeres
IT
        (-specific probes, used for mapping and estimating gene amplification;
        HER-2/neu gene amplification and expression
        in breast cancer)
     Gene, animal
TT
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (ERBB2; HER-2/neu gene amplification and
        expression in breast cancer)
TΥ
     Human
     Mammary gland, neoplasm
        (HER-2/neu gene amplification and
        expression in breast cancer)
IT
     neu (receptor)
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (HER-2/neu gene amplification and
        expression in breast cancer)
     Probes (nucleic acid)
TT
     RL: BUU (Biological use, unclassified); BIOL (Biological study); USES
     (Uses)
         (HER-2/new gene amplification and
        expression in breast cancer)
     Chromosome
TT
        (human 17, chromosomal location of HER-2/
        neu gene; HER-2/neu gene
        amplification and expression in breast cancer)
IT
     Nucleic acid hybridization
         (in situ, fluorescence, for HER-2/neu
        gene; HER-2/neu gene amplification and
        expression in breast cancer)
IT
     Genetic mapping
         (of HER-2/neu gene; HER-
        2/neu gene amplification and expression in breast
        cancer)
IT
     Gene
         (processes, gene amplification; HER-2/neu
        gene amplification and expression in breast cancer)
```

REFERENCE COUNT: 58 THERE ARE 58 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L84 ANSWER 19 OF 40 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2002:279180 CAPLUS

DOCUMENT NUMBER: 136:291216

TITLE: Electrochemical characterization of screen-printed

carbonaceous electrodes for the determination of

peroxidase activity in novel screen-printed

flow-through modules

AUTHOR(S): Stiene, Matthias; Bilitewski, Ursula

CORPORATE SOURCE: German Research Centre for Biotechnology,

Braunschweig, 38124, Germany

SOURCE: Analytical and Bioanalytical Chemistry (2002), 372(2),

240-247

CODEN: ABCNBP; ISSN: 1618-2642

PUBLISHER: Springer-Verlag

DOCUMENT TYPE: Journal LANGUAGE: English

ED Entered STN: 15 Apr 2002

AB A novel totally screen-printed flow-through cell for immunoanal. is

presented. It contained screen-printed carbonaceous electrodes, which

allowed the determination of peroxidase activity through the electrochem.

reduction of

p-benzoquinone. As different electrode materials differ strongly in their electrochem. properties, electrodes resulting from various screen-printable carbonaceous pastes were characterized using the hydroquinone/p-benzoquinone redox couple. For most of the electrodes, cyclic voltammogram peak sepns. of between 550 and 670 mV were observed indicating only quasi-reversible electrochem. behavior. This was confirmed by variation of the peak separation with scan rate. Heterogeneous electron transfer rates of .apprx.0.5-1+10-3 cm s-1 and electrochem. activation energies of .apprx.20 kJ mol-1 were found. These flow-through cells were not only applied to electrochem. peroxidase activity detns. but also, in combination with a sep. detector, as affinity reactors. After biotinylation of screen-printed layers, streptavidin and then biotinylated peroxidase could be bound. However, as signals were only 10-20% of those obtained with a column filled with biotinylated glass beads, only the screen-printed electrochem. detector was applied to the detection of antibodies against the African Swine Fever Virus.

CC 9-7 (Biochemical Methods)

ST screen printed carbonaceous electrode detn peroxidase activity

IT Activation energy

Affinity

African swine fever virus

Biotinylation

Cyclic voltammetry

Electron transfer

Enzyme electrodes

Flow

Reduction, electrochemical

Screen printing

(electrochem. characterization of screen-printed carbonaceous electrodes for determination of peroxidase activity in novel screen-printed flow-through modules)

IT Antibodies and Immunoglobulins

RL: ANT (Analyte); ANST (Analytical study)

(electrochem. characterization of screen-printed carbonaceous electrodes for determination of **peroxidase** activity in novel screen-printed flow-through modules)

```
TΤ
     Glass beads
     RL: DEV (Device component use); USES (Uses)
        (electrochem. characterization of screen-printed carbonaceous
        electrodes for determination of peroxidase activity in novel
        screen-printed flow-through modules)
IT
     Electrodes
        (glassy carbon; electrochem. characterization of screen-printed
        carbonaceous electrodes for determination of peroxidase activity in
        novel screen-printed flow-through modules)
     106-51-4, p-Benzoquinone, analysis 7722-84-1, Hydrogen
IT
     peroxide, analysis 9003-99-0, Peroxidase
     RL: ANT (Analyte); ANST (Analytical study)
        (electrochem. characterization of screen-printed carbonaceous
        electrodes for determination of peroxidase activity in novel
        screen-printed flow-through modules)
     123-31-9, Hydroquinone, analysis
IT
     Streptavidin
     RL: ARU (Analytical role, unclassified); ANST (Analytical study)
        (electrochem. characterization of screen-printed carbonaceous
        electrodes for determination of peroxidase activity in novel
        screen-printed flow-through modules)
     1344-28-1, Aluminum oxide, uses
                                      7440-44-0, Carbon, uses
IT
     7440-57-5, Gold, uses 7782-42-5, Graphite, uses
     RL: DEV (Device component use); USES (Uses)
        (electrochem. characterization of screen-printed carbonaceous
        electrodes for determination of peroxidase activity in novel
        screen-printed flow-through modules)
REFERENCE COUNT:
                         42
                               THERE ARE 42 CITED REFERENCES AVAILABLE FOR THIS
                               RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
L84 ANSWER 20 OF 40 CAPLUS COPYRIGHT 2005 ACS on STN
                         2002:122051 CAPLUS
ACCESSION NUMBER:
DOCUMENT NUMBER:
                         137:243010
TITLE:
                         Fluorescence in situ hybridization analysis of
                         HER-2/neu in brushings of
                         normal oral mucosa
                         Paradiso, Angelo; Abatangelo, Marta; Piepoli, Sandra;
AUTHOR (S):
                         Tommasi, Stefania; Xu, Jian-Ming; Caponio, Maria
                         Angela; Marzullo, Franco; D'Auria, Carlo; Achille,
                         Gaetano; Grammatica, Luciano
CORPORATE SOURCE:
                         Clinical Experimental Oncology Laboratory, National
                         Cancer Institute, Bari, Italy
                         Cancer Genetics and Cytogenetics (2002), 132(2),
SOURCE:
                         141-144
                         CODEN: CGCYDF; ISSN: 0165-4608
                         Elsevier Science Inc.
PUBLISHER:
DOCUMENT TYPE:
                         Journal
                         English
LANGUAGE:
     Entered STN: 15 Feb 2002
ED
AB
     Oncogene alterations have been clearly demonstrated to be related to the
     carcinogenesis and progression of oral squamous cell carcinoma (OSCC).
     However, the anal. of these alterations for screening and early diagnostic
     purposes generally requires invasive techniques for surgical removal of
     pathol. epithelium. The aim of the present study was to assess the
     feasibility of fluorescence in situ hybridization (FISH) anal. of
     HER-2/neu amplification in oral mucosa
     brushings and to compare the HER-2/neu
     status with the history and smoking and drinking habits of healthy
     subjects. Cells obtained by centrifugation of oral brushings from 21
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subjects (overall number of cells: 5125) were suspended in physiol. saline

and fixed onto two slides for cytol. evaluation and FISH anal. (dual-target, dual-color fluorescence assay) of the HER-2/neu gene and CEP17 centromere. A mean of 89.8% of the cells showed two HER-2/neu signals and a mean of 94% had two CEP17 signals at fluorescent microscopy. Finally, a mean of 96% of cells with HER-2/neu / CEP17 had a ratio equal to 1. No association between smoking and drinking habits, age and the HER-2/neu and CEP17 characteristics evaluated by FISH was found. 3-6 (Biochemical Genetics) CC Section cross-reference(s): 9, 13 Probes (nucleic acid) IT RL: ARG (Analytical reagent use); BUU (Biological use, unclassified); ANST (Analytical study); BIOL (Biological study); USES (Uses) (CEP17; fluorescence in situ hybridization anal. of HER-2/neu in brushings of normal oral mucosa) IT Gene, animal RL: ANT (Analyte); BSU (Biological study, unclassified); ANST (Analytical study); BIOL (Biological study) (ERBB2; fluorescence in situ hybridization anal. of HER-2/neu in brushings of normal oral mucosa) ΙT Health Human (fluorescence in situ hybridization anal. of HER-2/ neu in brushings of normal oral mucosa) TΤ Chromosome (human 17, HER-2/neu gene maps to; fluorescence in situ hybridization anal. of HER-2/ neu in brushings of normal oral mucosa) Nucleic acid hybridization IT (in situ, fluorescence; fluorescence in situ hybridization anal. of HER-2/neu in brushings of normal oral mucosa) ·IT Mouth (mucosa, brushings o; fluorescence in situ hybridization anal. of HER-2/neu in brushings of normal oral mucosa) REFERENCE COUNT: THERE ARE 18 CITED REFERENCES AVAILABLE FOR THIS 18 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT L84 ANSWER 21 OF 40 CAPLUS COPYRIGHT 2005 ACS on STN ACCESSION NUMBER: 2001:64266 CAPLUS 134:97508 DOCUMENT NUMBER: Metal nanoshells for biosensing applications TITLE: West, Jennifer L.; Sershen, Scott R.; Halas, Nancy L.; INVENTOR(S): Oldenburg, Steven J.; Averitt, Richard D. PATENT ASSIGNEE(S): Wm. Marsh Rice University, USA SOURCE: PCT Int. Appl., 39 pp. CODEN: PIXXD2 DOCUMENT TYPE: Patent LANGUAGE: English FAMILY ACC. NUM. COUNT: PATENT INFORMATION: PATENT NO. KIND DATE APPLICATION NO. DATE

	***			2
WO 2001006257	A1	20010125	WO 2000-US19375	20000714
W: CA, JP RW: AT. BE. CH.	CY. DE	. DK. ES.	FI, FR, GB, GR, IE, IT,	LU. MC. NI.
PT, SE	,	,,,	11, 11, 12, 11, 12, 11,	

0

CA 2377722 AA 20010125 CA 2000-2377722 20000714 20020605 EP 2000-952155 EP 1210600 A1 20000714 AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, FI, CY 20030204 JP 2001-510843 JP 2003504642 T2 20000714 US 1999-144136P PRIORITY APPLN. INFO.: P 19990716 WO 2000-US19375 W 20000714 Entered STN: 26 Jan 2001 ED The present invention provides nanoshell particles ("nanoshells") for use AB in biosensing applications, along with their manner of making and methods of using the nanoshells for in vitro and in vivo detection of chemical and biol. analytes, preferably by surface enhanced Raman light scattering. The preferred particles have a non-conducting core and a metal shell surrounding the core. For given core and shell materials, the ratio of the thickness (i.e., radius) of the core to the thickness of the metal shell is determinative of the wavelength of maximum absorbance of the particle. By controlling the relative core and shell thicknesses, biosensing metal nanoshells are fabricated which absorb light at any desired wavelength across the UV to IR range of the electromagnetic spectrum. The surface of the particles are capable of inducing an enhanced SERS signal that is characteristic of an analyte of interest. certain embodiments a biomol. is conjugated to the metal shell and the SERS signal of a conformational change or a reaction product is detected. IC ICM G01N033-543 G01N033-566; G01N033-551; G01N033-553; G01N015-06; G01N033-00; G01N033-48; G01N027-02; G01N027-06; G01N027-12 9-5 (Biochemical Methods) CC IT Antibodies Antigens Enzymes, uses Metals, uses Oligonucleotides Polysaccharides, uses Proteins, specific or class RL: ARG (Analytical reagent use); TEM (Technical or engineered material use); ANST (Analytical study); USES (Uses) (conjugates; metal nanoshells for biosensing applications) IT Antibodies Antigens Oligonucleotides Peptides, analysis Polysaccharides, analysis Proteins, general, analysis RL: ANT (Analyte); ANST (Analytical study) (metal nanoshells for biosensing applications) REFERENCE COUNT: THERE ARE 10 CITED REFERENCES AVAILABLE FOR THIS 10 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT L84 ANSWER 22 OF 40 CAPLUS COPYRIGHT 2005 ACS on STN 2001:244546 CAPLUS ACCESSION NUMBER: DOCUMENT NUMBER: 135:3482 TITLE: Scavenging of extracellular H2O2 by catalase inhibits the proliferation of HER-2/ Neu-transformed Rat-1 fibroblasts through the induction of a stress response Preston, Thomas J.; Muller, William J.; Singh, Gurmit AUTHOR (S): Hamilton Regional Cancer Centre, Faculty of Health CORPORATE SOURCE: Sciences, McMaster University, Hamilton, ON, L8V 5C2, Can.

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Journal of Biological Chemistry (2001), 276(12),
SOURCE:
                          9558-9564
                          CODEN: JBCHA3; ISSN: 0021-9258
                          American Society for Biochemistry and Molecular
PUBLISHER:
                          Biology
DOCUMENT TYPE:
                          Journal
LANGUAGE:
                          English
     Entered STN: 06 Apr 2001
ED
AB
     High levels of reactive oxygen species (ROS) are associated with
     cytotoxicity. Alternatively, nontoxic levels of ROS like hydrogen
     peroxide (H2O2) can mediate the transmission of many intracellular
     signals, including those involved in growth and transformation. To
     identify pathways downstream of endogenous cellular H2O2 production, the
     response of Rat-1 fibroblasts exhibiting differential HER-
     2/Neu receptor tyrosine kinase activity to removal of physiol. H2O2 concns. was investigated. The proliferation of all cells
     was abolished by addition of the H2O2 scavenger catalase to the culture
     medium. HER-2/Neu activity was not
     significantly affected by catalase treatment, suggesting that the
     target(s) of the H2O2 signal lie downstream of the receptor in our model.
     ERK1/2 phosphorylation was blocked by catalase in fibroblasts expressing
     wild type Neu, however such a response did not occur in cells possessing
     activated mutant Neu. This indicates that the ERK1/2 response contributes
     little to the growth inhibition observed By contrast, JNK1 activity
     increased following the addition of catalase or H2O2, regardless of Neu
     activity or level of cell transformation. Phosphorylation of p38 MAPK was
     induced by H2O2 but not by catalase. These observations suggest that scavenging of H2O2 from the cellular environment blocks Rat-1
     proliferation primarily through the activation of stress pathways.
CC
     13-6 (Mammalian Biochemistry)
ST
     fibroblast proliferation hydrogen peroxide catalase
     Neu phosphorylation oxidn apoptosis
IT
     Phosphorylation, biological
        (protein; scavenging of extracellular H2O2 by catalase inhibits the
        proliferation of HER-2/Neu-transformed
        Rat-1 fibroblasts through the induction of a stress response)
IT
     Apoptosis
     Cell proliferation
     Fibroblast
     Oxidative stress, biological
     Signal transduction, biological
        (scavenging of extracellular H2O2 by catalase inhibits the
        proliferation of HER-2/Neu-transformed
        Rat-1 fibroblasts through the induction of a stress response)
IT
     9001-05-2, Catalase
     RL: BAC (Biological activity or effector, except adverse); BSU (Biological
     study, unclassified); BIOL (Biological study)
        (scavenging of extracellular H2O2 by catalase inhibits the
        proliferation of HER-2/Neu-transformed
        Rat-1 fibroblasts through the induction of a stress response)
IT
     7722-84-1, Hydrogen peroxide, biological
     studies
               137632-09-8
                              142243-02-5D, Mitogen-activated protein kinase,
     8 Eq
     RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL
     (Biological study); PROC (Process)
        (scavenging of extracellular H2O2 by catalase inhibits the
        proliferation of HER-2/Neu-transformed
        Rat-1 fibroblasts through the induction of a stress response)
REFERENCE COUNT:
                                THERE ARE 39 CITED REFERENCES AVAILABLE FOR THIS
                          39
                                RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
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L84 ANSWER 23 OF 40 CAPLUS COPYRIGHT 2005 ACS on STN 2000:729505 CAPLUS ACCESSION NUMBER: 135:56565 DOCUMENT NUMBER: HER-2/neu oncogene TITLE: amplification determined by fluorescence in situ hybridization Ross, Jeffrey S.; Sheehan, Christine E.; Fletcher, AUTHOR (S): Jonathan A. Department of Pathology and Laboratory Medicine, CORPORATE SOURCE: Albany Medical College, Albany, NY, USA Methods in Molecular Medicine (2001), 49, 93-104 SOURCE: CODEN: MMMEFN Humana Press Inc. PUBLISHER: DOCUMENT TYPE: Journal English LANGUAGE: Entered STN: 16 Oct 2000 ED The proto-oncogene HER-2/neu (C-erbB-2) has AB been localized to chromosome 17q and encodes a transmembrane tyrosine kinase growth factor receptor. In addition to its association with disease outcome in gastrointestinal, pulmonary, genitourinary, and other neoplasms, amplification of the C-erbB-2 gene or over-expression of its protein has been identified in breast cancers. In this chapter, we describe the use of a fluorescence in situ hybridization (FISH) assay, for determining HER-2/neu (C-erbB-2) oncogene amplification in breast cancer, which is based on the Oncor INFORM® HER-2/neu Gene Detection System, and which uses a biotinylated probe. CC 3-1 (Biochemical Genetics) Section cross-reference(s): 14 Recombination, genetic IT(amplification; HER-2/neu oncogene amplification determined by fluorescence in situ hybridization) Probes (nucleic acid) IT RL: ARG (Analytical reagent use); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); USES (Uses) (biotinylated; HER-2/neu oncogene amplification determined by fluorescence in situ hybridization) IT Gene, animal RL: ADV (Adverse effect, including toxicity); ANT (Analyte); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); USES (Uses) (c-erbB2; HER-2/neu oncogene amplification determined by fluorescence in situ hybridization) Nucleic acid hybridization TT (in situ, fluorescence; HER-2/neu oncogene amplification determined by fluorescence in situ hybridization) Diagnosis IT (mol.; HER-2/new oncogene amplification determined by fluorescence in situ hybridization) Mammary gland TΤ (neoplasm; HER-2/neu oncogene amplification determined by fluorescence in situ hybridization) REFERENCE COUNT: 56 THERE ARE 56 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT L84 ANSWER 24 OF 40 CAPLUS COPYRIGHT 2005 ACS on STN 2000:900853 CAPLUS ACCESSION NUMBER: DOCUMENT NUMBER: 134:39172 Markers for prostate cancer TITLE:

INVENTOR(S):
PATENT ASSIGNEE(S):

Cordon-Cardo, Carlos; Scher, Howard I.; Koff, Andrew Sloan-Kettering Institute for Cancer Research, USA

PCT Int. Appl., 128 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

SOURCE:

Patent English

LANGUAGE:

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

P	ATENT	NO.			KINI	D	DATE			APF	LL	CAT	ION	. O <i>n</i>		D	ATE	
-						_										-		
W	O 2000	0772	58		A1		2000	1221	1	WO	20	00-1	US16	007		2	0000	609
	W:	CA,	JP,	US														
	RW:	ΑT,	BE,	CH,	CY,	DE,	DK,	ES,	FI,	FF	۷, ۱	GB,	GR,	ΙE,	IT,	LU,	MC,	NL,
		PT,	SE															
C	A 2375	228			AA		2000	1221		CA	20	00-2	2375	228		2	0000	609
E	P 1208	232			A1		2002	0529		ΕP	20	00-	9382	56		2	0000	609
	R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GF	₹,	IT,	LI,	LU,	NL,	SE,	MC,	PT,
		IE,	FI,	CY														
E	P 1512	755			A2		2005	0309		ΕP	20	04-2	2300	0		2	0000	609
	R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GF	₹,	IT,	LI,	LU,	NL,	SE,	MC,	PT,
		IE,	FI,	CY														
PRIORI'	TY APP	LN.	INFO	. :					1	US	19	99-3	3299	17		A2 1	9990	610
										ΕP	20	00-	9382	56		A3 2	0000	609
									1	WO	20	00-1	US16	007		W 2	0000	609

ED Entered STN: 22 Dec 2000

- This invention provides a method for determining the aggressiveness of a AB prostate carcinoma comprising: (a) obtaining a sample of the prostate carcinoma; and (b) detecting the presence of p27 protein in the prostate carcinoma, the absence of p27 indicating that the prostate carcinoma is aggressive. This invention also provides a method for diagnosing a benign prostate hyperplasia comprising: (a) obtaining an appropriate sample of the hyperplasia; and (b) detecting the presence of the p27 RNA, a decrease of the p27 RNA indicating that the hyperplasia is benign. This invention provides various uses of p27 in prostate cancer. Finally, this invention also provides different marker for prostate cancer. To determine whether loss of p27 expression was a common feature in prostate cancer, 74 prostate carcinomas from primary and metastatic sites, representing different hormone sensitivities were studied by immunohistochem. staining and in situ hybridization. Other markers such as cyclin D1, cyclin-dependent kinase inhibitor p16, and Her-2/neu were also studied.
- IC ICM C12Q001-68

ICS C07H021-04

CC 9-10 (Biochemical Methods)

Section cross-reference(s): 3, 14, 63

IT Antibodies

Probes (nucleic acid)

RL: ARG (Analytical reagent use); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); USES (Uses)

(markers for prostate cancer)

IT Androgens

RL: MSC (Miscellaneous)

(prostate cancer dependent on, treatment with antibody to Her -2/neu; markers for prostate cancer)

IT 1672-46-4D, Digoxigenin, conjugates with nucleic acid
probes

RL: ARG (Analytical reagent use); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); USES (Uses) (markers for prostate cancer)

THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: 3 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L84 ANSWER 25 OF 40 CAPLUS COPYRIGHT 2005 ACS on STN

2000:277876 CAPLUS ACCESSION NUMBER:

132:313678 DOCUMENT NUMBER:

Metal salt particle-adsorbed adjuvant systems and TITLE:

vaccines

Garcon, Nathalie INVENTOR(S):

Smithkline Beecham Biologicals SA, Belg. PATENT ASSIGNEE(S):

PCT Int. Appl., 37 pp. SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE: Patent English LANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

F	raq	ENT I	.00			KIN	D	DATE			APP	LICA	CION	NO.			DATE	
																	 19991	008
W	10	20000	0231	05		A3		2000	0803									
		W:	•			•		•									, CR,	
																	, ID,	
			IN,	IS,	JP,	KE,	KG,	ΚP,	KR,	ΚZ,	LC	LK,	LR,	LS,	LT,	LU	, LV,	MA,
			MD,	MG,	MK,	MN,	MW,	MX,	NO,	NZ,	PL	, PT	RO,	RU,	SD,	SE	, SG,	SI,
			SK,	SL,	ΤJ,	TM,	TR,	TT,	TZ,	UA,	UG	, US	UZ,	VN,	ΥU,	ZA	, ZW,	AM,
			AZ,	BY,	KG,	ΚZ,	MD,	RU,	ТJ,	TM								
		RW:	GH,	GM,	KE,	LS,	MW,	SD,	SL,	SZ,	TZ	, UG	ZW,	ΑT,	BE,	CH	, CY,	DE,
			DK,	ES,	FI,	FR,	GB,	GR,	ΙE,	IT,	LU	, MC	NL,	PT,	SE,	BF	, вJ,	CF,
			CG,	CI,	CM,	GA,	GN,	GW,	ML,	MR,	NE	, SN	TD,	TG				
C	'A	23470	099			AA		2000	0427		CA	1999	-2347	099			19991	800
Е	3R	9915	545			Α		2001	0814		BR	1999	-1554	5			19991	800
E	P	11268	376			A2		2001	0829		ΕP	1999	9706	07			19991	800
		R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR	, IT	LI,	LU,	NL,	SE	, MC,	PT,
			IE,	SI,	LT,	LV,	FI,	RO	•	-			-					
Т	'R	20010	0105	5		T2		2001	0921		TR	2001	-2001	0105	5		19991	800
A	U	75058	37			В2		2002	0725		AU	2000-	1151	8			19991	800
																	19991	
J	ΓP	20035	5190	84		Т2		2003	0617		JΡ	2000-	-5768	78			19991	800
E	P	1588	714			A2		2005	1026		ΕP	2005	7636	8			19991	800
		R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR	, IT	LI,	LU,	NL,	SE	, MC,	PT,
			IE,	SI,	FI,	CY												
Т	w	58693						2004	0511		TW	1999	8811	7873			19991	015
		20010									NO	2001	-1801				20010	409
Z	A	2001	0029	54		Α		2002	0520		zA	2001	2954				20010	410
PRIORI	TY	APPI	LN.	INFO	. :						GB	1998	-2270	3		Α	19981	016
																	19981	
																	19981	
											ΕP	1999-	9706	07		A 3	19991	800
											WO	1999-	EP77	64		W	19991	800
ED E	nt	ered	STN	: 28	8 Ap:	r 20	00											

28 Apr 2000

The present invention provides vaccine and adjuvant formulations AΒ comprising an immunostimulant and a metal salt. The immunostimulant is adsorbed onto a particle of metal salt (e.g. aluminum hydroxide or phosphate) and the resulting particle is essentially devoid of antigen.

IC ICM A61K039-39

ICS A61K039-00; A61K039-29; A61K039-015; A61P031-12; A61P031-04; A61P033-06; A61P035-00; A61P037-08

63-5 (Pharmaceuticals) CC

Section cross-reference(s): 15

IT Antigens

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); PEP (Physical, engineering or chemical process); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PROC (Process); USES (Uses)

(Her-2 neu; metal salt particle-adsorbed adjuvant systems and vaccines)

IT 7429-90-5D, Aluminum, salts, biological studies 7439-89-6D, Iron, salts, biological studies 7440-41-7D, Beryllium, salts, biological studies 7440-45-1D, Cerium, salts, biological studies 7440-47-3D, Chromium, salts, biological studies 7440-66-6D, Zinc, salts, biological studies 7440-70-2D, Calcium, salts, biological studies 7784-30-7, Aluminum phosphate 21645-51-2, Aluminum hydroxide, biological studies RL: PEP (Physical, engineering or chemical process); THU (Therapeutic use); BIOL (Biological study); PROC (Process); USES (Uses)

(metal salt particle-adsorbed adjuvant systems and vaccines)

L84 ANSWER 26 OF 40 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1999:435585 CAPLUS

DOCUMENT NUMBER: 131:56122

TITLE: Metal oxide matrix biosensors

INVENTOR(S): Rauh, R. David

PATENT ASSIGNEE(S): EIC Laboratories, Inc., USA

SOURCE: U.S., 17 pp.
CODEN: USXXAM

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 5922183	Α	19990713	US 1997-880615	19970623
PRIORITY APPLN. INFO.:			US 1997-880615	19970623

ED Entered STN: 15 Jul 1999

AB A thin film matrix for biomols., suitable for forming electrochem. and biosensors comprising a general class of materials known as hydrous metal oxides which are also conductive or semiconductive of electrons and which have been shown to have excellent stability against dissoln. or irreversible reaction in aqueous and nonaq. solns. The composites are bifunctional, providing both amperometric and potentiometric (pH) transduction. The thin film composites of the oxides and biol. mols. such as enzymes, antibodies, antigens and DNA strands can be used for both amperometric and potentiometric sensing. Hydrous Ir oxide is the preferred matrix embodiment, but conducting or semiconducting oxides, of Ru, Pd, Pt, Zr, Ti and Rh and mixts. thereof have similar features. The hydrous oxides are very stable against oxidation damage.

IC ICM G01N027-26

INCL 204403000

CC 9-1 (Biochemical Methods)

IT Antibodies

Antigens

DNA

Enzymes, uses

Oxides (inorganic), uses

RL: DEV (Device component use); USES (Uses)

(metal oxide matrix biosensors)

REFERENCE COUNT: 36 THERE ARE 36 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

```
L84 ANSWER 27 OF 40 CAPLUS COPYRIGHT 2005 ACS on STN
                         1999:236198 CAPLUS
ACCESSION NUMBER:
                         131:179403
DOCUMENT NUMBER:
                         Influence of chemotherapy on the expression of p53,
TITLE:
                         HER-2/neu and
                         proliferation markers in ovarian cancer
                         Nijman, Hans W.; Kenemans, Peter; Poort-Keesom, Ria J.
AUTHOR (S):
                         J.; Verstraeten, Rob A.; Mensdorff-Pouilly, Sylvia;
                         Verheijen, Rene H. M.; Melief, Cornelis J. M.;
                         Hilgers, Jo; Meijer, Chris J. L. M.
                         Department of Obstetrics and Gynaecology, Academic
CORPORATE SOURCE:
                         Hospital, Vrije Universiteit Amsterdam, Amsterdam,
                         Neth.
                         European Journal of Obstetrics & Gynecology and
SOURCE:
                         Reproductive Biology (1999), 83(2), 201-206
CODEN: EOGRAL; ISSN: 0301-2115
                         Elsevier Science Ireland Ltd.
PUBLISHER:
                         Journal
DOCUMENT TYPE:
                         English
LANGUAGE:
     Entered STN: 16 Apr 1999
     Objective: Mutated p53 and HER-2/neu play a
AΒ
     role in the etiol. of ovarian cancer. It is important to know whether the
     expression of these proteins is affected by platinum-containing chemotherapy.
     Study design: Together with the cell proliferation markers Ki-67 and PCNA,
     the expression of p53 and HER-2/neu was
     assessed before and after chemotherapy.
                                               Paraffin-embedded tumor sections
     from 20 patients with ovarian cancer and four patients with benign
     disorders of the ovaries (controls) were analyzed. The expression of p53
     was determined by the antibodies DO-1 and BP53-12. In addition to HER-
     2/neu and PCNA specific antibodies, MIB-1 was used to
     detect Ki-67. Results: The expression of all markers was higher in
     ovarian cancer patients than in non-malignant controls. MIB-1 showed a
     significant increase of expression after chemotherapy.
     2/neu, p53 and PCNA also showed a clear increase after
     treatment, but this was not statistically significant.
                                                              HER-
     2/neu is of prognostic relevance with respect to the
     response to chemotherapy and survival. Conclusion: The different markers
     tested all increase after chemotherapy, but the differences are not
     statistically significant. Low HER-2/neu
     expression correlates with good outcome at second look.
CC
     1-6 (Pharmacology)
     Section cross-reference(s): 14
IT
     Ovary, neoplasm
        (influence of platinum-containing chemotherapy on expression of p53,
        HER-2/neu and proliferation markers in
        ovarian cancer)
     Ki-67 antigen
TT
     Proliferating cell nuclear antigen
     RL: BAC (Biological activity or effector, except adverse); BSU (Biological
     study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES
     (Uses)
        (influence of platinum-containing chemotherapy on expression of p53,
        HER-2/neu and proliferation markers in
        ovarian cancer)
     neu (receptor)
IT
     p53 (protein)
     RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL
     (Biological study); PROC (Process)
        (influence of platinum-containing chemotherapy on expression of p53,
        HER-2/neu and proliferation markers in
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Ralph Gitomer 10/658,609 ovarian cancer) Ovary, neoplasm IT Ovary, neoplasm (inhibitors; influence of platinum-containing chemotherapy on expression of p53, HER-2/neu and proliferation markers in ovarian cancer) Antitumor agents IT Antitumor agents (ovary; influence of platinum-containing chemotherapy on expression of p53, HER-2/neu and proliferation markers in ovarian cancer) 7440-06-4D, Platinum, compds., biological studies IT RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses) (influence of platinum-containing chemotherapy on expression of p53, HER-2/neu and proliferation markers in ovarian cancer) THERE ARE 28 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: 28 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT L84 ANSWER 28 OF 40 CAPLUS COPYRIGHT 2005 ACS on STN ACCESSION NUMBER: 1999:52824 CAPLUS 130:263969 DOCUMENT NUMBER: Artificial metalloenzymes based on protein cavities: TITLE: exploring the effect of altering the metal ligand attachment position by site directed mutagenesis Davies, Ronald R.; Kuang, Hao; Qi, Dongfeng; Mazhary, AUTHOR (S): Aram; Mayaan, Evelyn; Distefano, Mark D. Department of Chemistry, University of Minnesota, CORPORATE SOURCE: Minneapolis, MN, 55455, USA Bioorganic & Medicinal Chemistry Letters (1999), 9(1), SOURCE: 79-84 CODEN: BMCLE8; ISSN: 0960-894X Elsevier Science Ltd. PUBLISHER: Journal DOCUMENT TYPE: English LANGUAGE: Entered STN: 26 Jan 1999 EDIn an effort to construct catalysts with enzyme-like properties, we are AB employing a small, cavity-containing protein as a scaffold for the attachment of catalytic groups. In earlier work we demonstrated that a phenanthroline ligand could be introduced into the cavity of the protein ALBP (adipocyte lipid binding protein) and used to catalyze ester hydrolysis. To examine the effect of positioning the phenanthroline catalyst at different locations within the protein cavity of a related protein, IFABP (intestinal fatty acid-binding protein), three new constructs - Phen60, Phen72 and Phen104 - were prepared Each new conjugate was characterized by UV/vis spectroscopy, fluorescence spectroscopy, guanidine hydrochloride denaturation, gel filtration chromatog., and CD spectroscopy to confirm the preparation of the desired construct. Anal. of reactions containing Ala-OiPr showed that Phen60 catalyzed ester hydrolysis

CC 7-5 (Enzymes)

IT Enzyme functional sites

(active; exploring effect of altering metal ligand

of attachment of the metal ligand within the protein cavity.

with less selectivity than ALBP-Phen while Phen72 promoted this same reaction with higher selectivity. Reactions with Tyr-OMe were catalyzed with higher selectivity by Phen60 and more rapidly by Phen104. These results demonstrate that both the rates and selectivities of hydrolysis reactions catalyzed by these constructs are dependent on the precise site

attachment position by site directed mutagenesis artificial metalloenzymes based on fatty acid-binding protein)

IT Enzymes, preparation

RL: BPN (Biosynthetic preparation); CAT (Catalyst use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(synthetic; exploring effect of altering metal

ligand attachment position by site directed mutagenesis artificial metalloenzymes based on fatty acid-binding protein)

7440-50-8DP, Copper, complex with phenanthroline-IFABP (intestinal fatty acid-binding protein) mutants, biological studies RL: BAC (Biological activity or effector, except adverse); BPN (Biosynthetic preparation); BSU (Biological study, unclassified); CAT (Catalyst use); PRP (Properties); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation and metalation of I-FABP (intestinal fatty acid-binding protein)-phenanthroline conjugates)

REFERENCE COUNT:

THERE ARE 38 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L84 ANSWER 29 OF 40 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1998:682565 CAPLUS

DOCUMENT NUMBER: 129:299013

TITLE: Assessing prostate cancer based on measuring

amplification of the HER-2/

neu/c-erbB2 gene

INVENTOR(S): Ross, Jeffrey S.; Muraca, Patrick J.

PATENT ASSIGNEE(S): Albany Medical College, USA

SOURCE: PCT Int. Appl., 33 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 3

PATENT INFORMATION:

	PAT	CENT I	. 00			KINI)	DATE		1	APPL	ICAT	ION I	. 01		D	ATE	
							-									-		-
	WO	98454	479			A1		1998	1015	1	WO 1	998-1	US66:	21		1	9980	403
		W:	ΑL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	CA,	CH,	CN,	CU,	CZ,	DE,
			DK,	EE,	ES,	FI,	GB,	GE,	GH,	HU,	IL,	IS,	JP,	KΕ,	KG,	ΚP,	KR,	ΚZ,
			LC,	LK,	LR,	LS,	LT,	LU,	LV,	MD,	MG,	MK,	MN,	MW,	MX,	NO,	NZ,	PL,
			PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	ΤJ,	TM,	TR,	TT,	UA,	UG,	UZ,
			VN,	YU,	ZW,	AM,	ΑZ,	BY,	KG,	ΚZ,	MD,	RU,	ТJ,	TM				
		RW:						SD,							CY,	DE,	DK,	ES,
			FI,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE,	BF,	ВJ,	CF,	CG,	CI,
			CM,	GA,	GN,	ML,	MR,	NE,	SN,	TD,	TG							
	US	5994	071			Α		1999	1130	1	US 1	997-	8327	45		1	9970	404
	CA	2285	929			AA		1998	1015		CA 1	998-	2285	929		1	9980	403
	AU	9869	494			A1		1998	1030		AU 1	998-	6949	4		1	9980	403
	ΕP	9758	03			A1		2000	0202		EP 1	998-	9152	67		1	9980	403
		R:	CH,	DE,	FR,	GB,	LI											
	US	2002						2002	0620	•	US 1	998-	1529	34		1	9980	914
PRIO	RIT	APP	LN.	INFO	. :					•	US 1	997-	8327	45		A 1	9970	404
										1	WO 1	998-	US66:	21	1	W 1	9980	403
											US 1	998-	8841	7		A2 1	9980	601

ED Entered STN: 28 Oct 1998

AB A method of determining the severity of prostatic cancer includes measuring the level of amplification of the HER-2/neu gene in a sample of prostate tissue by fluorescence in-situ hybridization and comparing the measured level of amplification of the HER-2/neu gene in the sample with the level of HER

```
-2/neu gene in normal prostate tissue. A method for
determining treatment for a patient afflicted with prostate cancer includes
determining whether the number of copies of HER-2/neu
gene in prostate cells from the patient exceeds four by using fluorescence
in-situ hybridization and aggressively treating such patient having
prostate cells with five or more copies of the HER-2/
neu gene.
ICM C12Q001-68
ICS C12P019-34; C07H021-02; C07H021-04
3-1 (Biochemical Genetics)
Section cross-reference(s): 9, 15
Recombination, genetic
   (amplification; assessing prostate cancer based on measuring
   amplification of the HER-2/neu/c-erbB2
   gene)
Gene dosage
Immunoassay
Prognosis
   (assessing prostate cancer based on measuring amplification of the
   HER-2/neu/c-erbB2 gene)
Gene, animal
RL: ADV (Adverse effect, including toxicity); ANT (Analyte); ANST
(Analytical study); BIOL (Biological study)
   (c-erbB2; assessing prostate cancer based on measuring amplification of
   the HER-2/neu/c-erbB2 gene)
Diagnosis
   (cancer; assessing prostate cancer based on measuring amplification of
   the HER-2/neu/c-erbB2 gene)
Neoplasm
   (diagnosis; assessing prostate cancer based on measuring amplification
   of the HER-2/neu/c-erbB2 gene)
Probes (nucleic acid)
RL: ARG (Analytical reagent use); THU (Therapeutic use); ANST (Analytical
study); BIOL (Biological study); USES (Uses)
   (digoxigenin or biotin-labeled; assessing prostate cancer based on
   measuring amplification of the HER-2/neu
   /c-erbB2 gene)
Antibodies
RL: ARG (Analytical reagent use); THU (Therapeutic use); ANST (Analytical
study); BIOL (Biological study); USES (Uses)
   (fluorophore-labeled; assessing prostate cancer based on measuring
   amplification of the HER-2/neu/c-erbB2
   gene)
Diagnosis
   (genetic; assessing prostate cancer based on measuring amplification of
   the HER-2/neu/c-erbB2 gene)
Proteins, specific or class
RL: ARG (Analytical reagent use); THU (Therapeutic use); ANST (Analytical
study); BIOL (Biological study); USES (Uses)
   (green fluorescent, antibody labeled with; assessing prostate cancer
   based on measuring amplification of the HER-2/
   neu/c-erbB2 gene)
Nucleic acid hybridization
   (in situ, fluorescence; assessing prostate cancer based on measuring
   amplification of the HER-2/neu/c-erbB2
   gene)
Prostate gland
   (neoplasm; assessing prostate cancer based on measuring amplification
   of the HER-2/neu/c-erbB2 gene)
58-85-5D, Biotin, DNA probe labeled with 1672-46-4D, Digoxigenin, DNA
```

IC

CC

IT

2321-07-5D, Fluorescein, antibody labeled with probe labeled with 70281-37-7D, Tetramethylrhodamine, antibody labeled with 82354-19-6D, Texas Red, antibody labeled with 117939-97-6D, antibody labeled with 146368-15-2D, Cy 5, antibody labeled with 146397-20-8D, Cy3, antibody labeled with

RL: ARG (Analytical reagent use); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); USES (Uses)

(assessing prostate cancer based on measuring amplification of the HER-2/neu/c-erbB2 gene)

THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: 6

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L84 ANSWER 30 OF 40 CAPLUS COPYRIGHT 2005 ACS on STN

1994:428608 CAPLUS ACCESSION NUMBER:

121:28608 DOCUMENT NUMBER:

Methods for the detection of chromosome structural TITLE:

abnormalities by (in situ) hybridization to fixed

INVENTOR (S): Wang, Mary G.; George, Albert L., Jr.; Light,

Elizabeth S.

Oncor, Inc., USA PATENT ASSIGNEE(S):

PCT Int. Appl., 51 pp. SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 9409022	A1	19940428	WO 1993-US9658	19931008
W: AU, CA, JP,	KR			
RW: AT, BE, CH,	DE, DK	, ES, FR, GB	B, GR, IE, IT, LU, MC,	NL, PT, SE
AU 9453555	A1	19940509	AU 1994-53555	19931008
US 5856089	Α	19990105	US 1994-279315	19940722
PRIORITY APPLN. INFO.:			US 1992~958907	A 19921009
			WO 1993-US9658	W 19931008

Entered STN: 23 Jul 1994 ED

- The present invention is directed to in situ hybridization methods using AB nucleic acid probes for single copy sequences for detecting chromosomal structural abnormalities in fixed tissue obtained from a patient suspected of having a chromosomal structural abnormality. The method comprises (1) obtaining a fixed tissue sample from patient, (2) digesting the fixed sample with an effective amount of proteinase, (3) performing in situ hybridization, and (4) comparing with a normal control and detecting chromosome structural abnormality. With the claimed method, neuroblastoma and bladder carcinoma were detected with N-myc and chromosome 15 probe, and breast carcinoma was determined with HER-2/neu probe.
- ICM C07H021-02 IC
 - ICS C07H021-04; C12Q001-68; C12P019-34
- 3-1 (Biochemical Genetics) CC
- 9001-78-9D, Alkaline phosphatase, conjugates with avidin or antibody IT 9003-99-0, Peroxidase

RL: USES (Uses)

(probe conjugates binding to, for chromosome structural abnormality determination by DNA hybridization)

L84 ANSWER 31 OF 40 BIOSIS COPYRIGHT (c) 2005 The Thomson Corporation on

stn

ACCESSION NUMBER: 2004:229847 BIOSIS DOCUMENT NUMBER: PREV200400232855

TITLE: Purification of the plant alternative oxidase from Arum

maculatum: Measurement, stability and metal requirement.

AUTHOR(S): Afformation Afformation Source: Department

Affourtit, Charles [Reprint Author]; Moore, Anthony L. Department of Biochemistry, School of Life Sciences, University of Sussex, Falmer, Brighton, BN1 9QG, UK

C.Affourtit@sussex.ac.uk

SOURCE:

Biochimica et Biophysica Acta, (15 February 2004) Vol.

1608, No. 2-3, pp. 181-189. print.

ISSN: 0006-3002 (ISSN print).

DOCUMENT TYPE:

Article English

LANGUAGE: ENTRY DATE:

Entered STN: 28 Apr 2004

Last Updated on STN: 28 Apr 2004

We have purified plant alternative oxidase (AOX) protein from the spadices of thermogenic Arum maculatum (cuckoo pint) to virtual homogeneity. The obtained enzyme fraction exhibits a high specific activity, consuming on average 32 mumol oxygen min-1 mg-1, which is completely stable for at least 6 months when the sample is stored at -70degreeC. This exceptionally stable AOX activity is inhibited approximately 90% (I50apprx10 muM) by 8-hydroxyquinoline (8-OHQ) and also, although to a lesser extent, by other metal chelators such as o-phenanthroline, alpha, alpha'-dipyridyl and EDTA. When inhibited by 8-OHQ, AOX activity is fully restored upon addition of 1.2 mM ferric iron, but neither ferrous iron nor manganese has any effect, whilst zinc decreases activity even further. Furthermore, we have developed a spectrophotometric assay to measure AOX activity in an accurate manner, which will facilitate future steady state and transient kinetic studies. The reliability of this assay is evidenced by retained stability of AOX protein during the course of the reaction, reproducibility of the measured initial rates, an observed 2:1 duroquinol-oxygen stoichiometry and by the fact that, in absolute terms, the measured rates of duroquinone formation and duroquinol disappearance are identical.

IT Major Concepts

Enzymology (Biochemistry and Molecular Biophysics)

IT Chemicals & Biochemicals

enzymes; metals; plant alternative

oxidase: activities, analysis, functions, measurement, metal requirement, properties, purification, stability; proteins

IT Methods & Equipment

protein purification methods: laboratory techniques

IT Miscellaneous Descriptors

enzyme activation

ORGN Classifier

Araceae 25230

Super Taxa

Monocotyledones; Angiospermae; Spermatophyta; Plantae

Organism Name

Arum maculatum (species)

Taxa Notes

Angiosperms, Monocots, Plants, Spermatophytes, Vascular Plants

ORGN Classifier

Plantae 11000

Super Taxa

Plantae

Organism Name

plant (common) Taxa Notes Plants

ANSWER 32 OF 40 BIOSIS COPYRIGHT (c) 2005 The Thomson Corporation on L84

2004:80628 BIOSIS ACCESSION NUMBER: DOCUMENT NUMBER: PREV200400082517

Enzymatic deposition and alteration of metals. TITLE: Hainfeld, James F. [Inventor, Reprint Author] AUTHOR (S):

Shoreham, NY, USA CORPORATE SOURCE:

ASSIGNEE: Nanoprobes, Yaphank, NY, USA

PATENT INFORMATION: US 6670113 20031230

SOURCE: Official Gazette of the United States Patent and Trademark

Office Patents, (Dec 30 2003) Vol. 1277, No. 5. http://www.uspto.gov/web/menu/patdata.html. e-file.

ISSN: 0098-1133 (ISSN print).

DOCUMENT TYPE: Patent LANGUAGE: English

Entered STN: 4 Feb 2004 ENTRY DATE:

Last Updated on STN: 4 Feb 2004

Disclosed are methods and materials for utilizing enzymes to act on metal ions in solution so that the ions are reduced to metal. Additionally, disclosed is how to use enzymes to accumulate metal particles. The alteration of metal particles by enzymes interacting with the organic shell of the particles is also described. These methods enable a wide range of applications including sensitive detection of genes and proteins, use as probes for microscopy, nanofabrication, biosensors, and remediation.

ΙT Major Concepts

Methods and Techniques

Chemicals & Biochemicals IT

metal particles

Methods & Equipment IT

enzymatic deposition methods: laboratory techniques; enzymatic metal alteration methods: laboratory techniques

ANSWER 33 OF 40 BIOSIS COPYRIGHT (c) 2005 The Thomson Corporation on L84

STN

ACCESSION NUMBER: 2002:185462 BIOSIS DOCUMENT NUMBER: PREV200200185462 Her-2/neu oncogene TITLE:

amplification in clinically localised prostate cancer. Oxley, J. D. [Reprint author]; Winkler, M. H.; Gillatt, D. AUTHOR (S):

A.; Peat, D. S.

CORPORATE SOURCE: Department of Cellular Pathology, Southmead Hospital,

Westbury on Trym, Bristol, BS10 5NB, UK

jon@jon-oxley.freeserve.co.uk

SOURCE: Journal of Clinical Pathology (London), (February, 2002)

Vol. 55, No. 2, pp. 118-120. print.

CODEN: JCPAAK. ISSN: 0021-9746.

DOCUMENT TYPE: Article LANGUAGE: English

ENTRY DATE: Entered STN: 6 Mar 2002

Last Updated on STN: 6 Mar 2002

Aim: To examine the incidence of Her-2/neu

oncogene amplification in clinically localised prostate cancer using in situ hybridisation. Methods: One hundred and seventeen patients, who had undergone radical prostatectomy, were identified and in situ hybridisation

was performed on formalin fixed, paraffin wax embedded tissue using the Ouantum Appliquee probe for Her-2/ neu. The enzyme peroxidase was used to detect the probe because this enabled a permanent record to be kept. Tumours in which there were five or more signals in each nucleus in > 20% of the tumour cells were considered to have a significantly increased copy number. A serial section from these tumours was then hybridised with the chromosome 17alpha satellite probe. The ratio of the percentage of cells showing an increase in Her-2/neu copy number to the number showing polysomy for chromosome 17 was calculated. A ratio above 2 was considered amplified. Results: Biochemical recurrence occurred in 50 (43%) patients and 24 (21%) had clinical recurrence. In situ hybridisation for Her-2/ neu was accessible in 114 (97%) patients. A significant increase in copy number was present in two patients (1.75%), but chromosome 17 hybridisation showed that the increase was the result of polysomy rather than true amplification. Both these patients had a Gleason score of 7 and stage T3; they also had recurrent clinical disease with distal metastasis within two and 19 months. Conclusions: Increased Her-2 /neu oncogene copy number appears to be rare in clinically localised prostatic adenocarcinoma and is related to chromosome 17 polysomy rather than true amplification. As a result, it would not be a useful biomarker for identifying those patients who will have recurrences after radical prostatectomy. Major Concepts Molecular Genetics (Biochemistry and Molecular Biophysics); Tumor Biology Diseases localized prostate cancer: neoplastic disease, reproductive system disease/male, urologic disease Prostatic Neoplasms (MeSH) ORGN Classifier Hominidae 86215 Super Taxa Primates; Mammalia; Vertebrata; Chordata; Animalia Organism Name human: male, patient Taxa Notes Animals, Chordates, Humans, Mammals, Primates, Vertebrates GEN human Her-2-neu oncogene (Hominidae): localized tumor amplification, tumor development role L84 ANSWER 34 OF 40 BIOSIS COPYRIGHT (c) 2005 The Thomson Corporation on STN 2001:496098 BIOSIS ACCESSION NUMBER: PREV200100496098 DOCUMENT NUMBER: Chromogenic in situ hybridization analysis of HER TITLE: -2/neu status in breast carcinoma: Application in screening of patients for trastuzumab (Herceptin(R)) therapy. Kumamoto, Hiroyuki [Reprint author]; Sasano, Hironobu; AUTHOR (S): Taniguchi, Takahiro; Suzuki, Takashi; Moriya, Takuya; Ichinohasama, Ryo Division of Oral Pathology, Department of Oral Medicine and CORPORATE SOURCE: Bioregulation, Graduate School of Dentistry, Tohoku University, 4-1 Seiryo-machi, Aoba-ku, Sendai, 980-8575, kumamoto@mail.cc.tohoku.ac.jp Pathology International, (August, 2001) Vol. 51, No. 8, pp. SOURCE: 579-584. print.

IT

TT

ISSN: 1320-5463.

DOCUMENT TYPE: Article LANGUAGE: English

ENTRY DATE: Entered STN: 24 Oct 2001

Last Updated on STN: 23 Feb 2002

AB Evaluation of HER-2/neu status is important

in the management of patients with breast carcinoma, especially in determining the possible application of trastuzumab, a humanized anti-

HER-2/neu monoclonal antibody. Chromogenic in

situ hybridization (CISH) detection of the HER-2/

neu oncogene is a newly developed in situ hybridization method that utilizes a robust and unique-sequence DNA probe labeled with digoxygenin, and sequential incubations with antidigoxygenin

fluorescein, antifluorescein peroxidase, and diaminobenzidine.

In this study, we examined 20 archival specimens of human breast carcinoma using CISH, and we correlated findings with immunohistochemical findings for HER-2/neu. HER-2/

neu immunohistochemistry was carried out with HercepTestTM, a standardized immunohistochemical examination system for HER-

2/new overexpression in surgical pathology specimens.

CISH analysis could be done in 18 out of 20 cases examined. Gene copy signals for HER-2/neu were recognized as

intranuclear brown dots in both neoplastic and non-neoplastic cells. Seven carcinomas showed an increased number or size of signals and were interpreted as being positive for HER-2/neu

amplification. Eight cases were positive with the HercepTestTM. Seven out of eight carcinoma cases found to overexpress immunoreactive HER-2/neu also demonstrated HER-

2/neu gene amplification following CISH analysis. There was a significant correlation between immunohistochemical and CISH analyses (P<0.001). We found that CISH was a specific, sensitive and easily applicable method for the detection of HER-2/neu gene amplification, which may be used together with

immunohistochemical examination for the evaluation of patients with breast carcinoma.

IT Major Concepts

Molecular Genetics (Biochemistry and Molecular Biophysics); Oncology (Human Medicine, Medical Sciences); Pharmacology

IT Diseases

breast carcinoma: neoplastic disease, reproductive system disease/female, drug treatment screening, gene expression status Breast Neoplasms (MeSH); Carcinoma (MeSH)

IT Chemicals & Biochemicals

chromogenic in-situ hybridization: genetic method; trastuzumab [Herceptin]: antineoplastic-drug, patient screening

ORGN Classifier

Hominidae 86215

Super Taxa

Primates; Mammalia; Vertebrata; Chordata; Animalia Organism Name

human: female, patient

Taxa Notes

Animals, Chordates, Humans, Mammals, Primates, Vertebrates

RN 180288-69-1 (trastuzumab)

180288-69-1 (Herceptin)

GEN human HER-2-neu oncogene (Hominidae): drug

treatment screening application, tumor expression status analysis

L84 ANSWER 35 OF 40 BIOSIS COPYRIGHT (c) 2005 The Thomson Corporation on STN

ACCESSION NUMBER: 2002:591751 BIOSIS DOCUMENT NUMBER: PREV200200591751

TITLE: Interstitial lung disease induced by exogenous agents:

Factors governing susceptibility.

AUTHOR(S): Nemery, B. [Reprint author]; Bast, A.; Behr, J.; Borm, P.

J. A.; Bourke, S. J.; Camus, Ph.; De Vuyst, P.; Jansen, H. M.; Kinnula, V. L.; Lison, D.; Pelkonen, O.; Saltini, C.

CORPORATE SOURCE: Laboratorium voor Pneumologie (Longtoxicologie), K.U.

Leuven, Herestraat 49, B-3000, Leuven, Belgium

SOURCE: European Respiratory Journal, (September, 2001) Vol. 18,

No. Supplement 32, pp. 30s-42s. print.

CODEN: ERJOEI. ISSN: 0903-1936.

DOCUMENT TYPE: Article LANGUAGE: English

ENTRY DATE: Entered STN: 13 Nov 2002

Last Updated on STN: 13 Nov 2002

The purpose of this review is to describe the present state of knowledge regarding host susceptibility factors that may determine the occurrence, development and severity of interstitial lung disease (ILD) caused by exogenous agents. First, host susceptibility may pertain to differences in the delivery and/or persistence of the noxious agent in the lung. The deposition and clearance of inhaled particles or fibres may vary depending on innate anatomical or physiological characteristics, and on acquired changes, such as nasal disease or smoking-induced alterations. Genetically- or environmentally-induced interindividual differences in the expression of pulmonary biotransformation enzymes may form the basis for, or contribute to the risk of, drug-induced interstitial lung disease. Secondly, there are genetic and acquired variations in various enzymatic and nonenzymatic defence systems that protect cells and tissues against oxidative stress, which is often involved in the pathogenesis of interstitial lung disease caused by particles, fibres, metals, organic agents and drugs. Thirdly, the occurrence of immunological sensitization is dependent on both genetic and environmental factors. This has been demonstrated in chronic beryllium lung disease and in hypersensitivity pneumonitis. Fourthly, the propensity of individuals to develop particular types of inflammation, such as granulomas, is probably under genetic control. The regulation and resolution of inflammation and fibrogenesis caused by dust particles are also partly determined by genetic factors, involving cytokine networks and growth factors. In conclusion, although the issue of genetics pervades the entire discussion of host susceptibility, genes are not the only determinants of health and disease. Environmental factors may be equally important in shaping host susceptibility. Therefore, research must be focused on both the genetic bases and the environmental determinants of interstitial lung disease, in order to provide mechanism-based prevention strategies, early detection of, and improved therapy for these conditions.

IT Major Concepts

Pulmonary Medicine (Human Medicine, Medical Sciences); Toxicology

IT Parts, Structures, & Systems of Organisms

lung: respiratory system

IT Diseases

chronic beryllium lung disease: respiratory system disease, toxicity

IT Diseases

interstitial lung disease: respiratory system disease, toxicity, drug-induced, etiology

Lung Diseases, Interstitial (MeSH)

IT Diseases

nasal disease: respiratory system disease

IT Chemicals & Biochemicals

beryllium: toxin; cytokine; drug: pharmaceutical, toxicity; fiber:

toxicity; growth factor; inhaled particle: deposition; metal: toxin; organic agent: toxin; pulmonary biotransformation enzyme: expression Miscellaneous Descriptors TΤ disease susceptibility; oxidative stress; smoking ORGN Classifier Hominidae 86215 Super Taxa Primates; Mammalia; Vertebrata; Chordata; Animalia Organism Name human: patient Taxa Notes Animals, Chordates, Humans, Mammals, Primates, Vertebrates 7440-41-7 (beryllium) RNL84 ANSWER 36 OF 40 BIOSIS COPYRIGHT (c) 2005 The Thomson Corporation on STN ACCESSION NUMBER: 2001:70020 BIOSIS DOCUMENT NUMBER: PREV200100070020 Chromogenic in situ hybridization (CISH): A practical new TITLE: alternative to FISH to detect HER-2/ neu amplification in archival breast cancer samples. Tanner, M. M. [Reprint author]; Gancberg, D.; DiLeo, A.; AUTHOR (S): Larsimont, D.; Rouas, G.; Piccart, M.; Isola, J. J. [Reprint author] Institute of Medical Technology, Univ Tampere, Tampere, CORPORATE SOURCE: Finland Breast Cancer Research and Treatment, (November, 2000) Vol. SOURCE: 64, No. 1, pp. 100. print. Meeting Info.: 23rd Annual San Antonio Breast Cancer Symposium. San antonio, Texas, USA. December 06-09, 2000. Cancer Therapy and Research Center Research Foundation. CODEN: BCTRD6. ISSN: 0167-6806. Conference; (Meeting) DOCUMENT TYPE: Conference; Abstract; (Meeting Abstract) LANGUAGE: English Entered STN: 7 Feb 2001 ENTRY DATE: Last Updated on STN: 12 Feb 2002 Major Concepts IT Biochemistry and Molecular Biophysics; Molecular Genetics (Biochemistry and Molecular Biophysics); Methods and Techniques; Tumor Biology Parts, Structures, & Systems of Organisms IT breast: reproductive system; tumor tissue, archival formalin-fixed, paraffin-embedded IT Diseases breast cancer: neoplastic disease, reproductive system disease/female Breast Neoplasms (MeSH) Chemicals & Biochemicals IT CB-11 monoclonal antibody; HER-2/neu [human epidermal growth factor receptor-2/neu]; HER-2 /neu probe [human epidermal growth factor receptor-2/neu probe]: digoxigenin-labelled; anti-digoxygenin-fluorescein [anti-digoxigenin-fluorescein]; anti-fluorescein-peroxidase; diaminobenzidine; gene copy cluster: intranuclear, peroxidase-positive; trastuzumab [Herceptin]: antineoplastic-drug TΤ Methods & Equipment brightfield microscopy: microscopy method; chromogenic in situ hybridization [CISH]: bioassay method, histochemical method;

fluorescence in-situ hybridization [FISH]: bioassay method, histochemical method; p185 immunohistochemistry: bioassay method, histochemical method, immunologic method

Miscellaneous Descriptors

Meeting Abstract

ORGN Classifier

Hominidae 86215

Super Taxa

Primates; Mammalia; Vertebrata; Chordata; Animalia

Organism Name

human: patient

Taxa Notes

Animals, Chordates, Humans, Mammals, Primates, Vertebrates

66836-18-8 (diaminobenzidine) RN

180288-69-1 (trastuzumab)

180288-69-1 (Herceptin)

GEN HER-2/neu gene [human epidermal growth

factor receptor-2/neu gene] (Hominidae): amplification, oncogene

L84 ANSWER 37 OF 40 BIOSIS COPYRIGHT (c) 2005 The Thomson Corporation on

ACCESSION NUMBER: 1997:301993 BIOSIS DOCUMENT NUMBER: PREV199799601196

TITLE: Antioxidant responses to simulated acid rain and heavy

metal deposition in birch seedlings.

AUTHOR (S): Korcheva, Julia [Reprint author]; Roy, Sashwati; Vranjic,

John A.; Haukioja, Erkki [Reprint author]; Hughes, Patrick

R.; Hanninen, Osmo

CORPORATE SOURCE: Lab. Ecological Zool., Dep. Biol., Univ. Turku, FIN-20014

Turku, Finland

SOURCE : Environmental Pollution, (1997) Vol. 95, No. 2, pp.

249-258.

CODEN: ENPOEK. ISSN: 0269-7491.

DOCUMENT TYPE: Article LANGUAGE:

English

ENTRY DATE: Entered STN: 9 Jul 1997

Last Updated on STN: 9 Jul 1997

AB This study measured the responses of different antioxidants in 2-year-old birch (Betula pendula Roth) seedlings subjected to simulated acid rain (pH 4.0) and heavy metals (Cu/Ni), applied alone or in combination for 2 The applied concentrations of pollutants did not significantly affect seedling biomass or total glutathione levels. Acid rain alone increased superoxide dismutase (SOD) activity both in leaves and roots, while heavy metals alone inhibited SOD activity in roots. Both acid rain and heavy metals applied singly increased ascorbate peroxidase (APX) and guaiacol peroxidase (GPX) activities in leaves but decreased activities in In contrast, acid rain and heavy metal treatments increased glutathione reductase (GR) activity in roots but not in leaves. Spraying birch seedlings with a mixture of acid rain and heavy metals increased SOD, APX and GPX activities in leaves and GR activity in roots. However, the effects of mixed pollutants on enzyme activities usually were less than the summed effects of individual pollutants. Enzyme responses also depended on where pollutants were applied: spraying pollutants onto the shoots initiated higher responses in SOD, APX and GPX than did application to the soil surface, while the opposite was true for GR.

ΤТ Major Concepts

> Biochemistry and Molecular Biophysics; Enzymology (Biochemistry and Molecular Biophysics); Pollution Assessment Control and Management Chemicals & Biochemicals

COPPER; NICKEL; GLUTATHIONE; ASCORBATE PEROXIDASE; GUAIACOL PEROXIDASE;

IT

GLUTATHIONE REDUCTASE Miscellaneous Descriptors TT ACID RAIN; ANTIOXIDANT; ASCORBATE PEROXIDASE; COPPER; DEPOSITION; ENZYMOLOGY; GLUTATHIONE; GLUTATHIONE REDUCTASE; GUAIACOL PEROXIDASE; HEAVY METAL; NICKEL; POLLUTION; SEEDLING ORGN Classifier Betulaceae 25645 Super Taxa Dicotyledones; Angiospermae; Spermatophyta; Plantae Organism Name birch Betula pendula Taxa Notes Angiosperms, Dicots, Plants, Spermatophytes, Vascular Plants RN7440-50-8 (COPPER) 7440-02-0 (NICKEL) 70-18-8 (GLUTATHIONE) 72906-87-7 (ASCORBATE PEROXIDASE) 9003-99-0 (GUAIACOL PEROXIDASE) 9001-48-3 (GLUTATHIONE REDUCTASE) ANSWER 38 OF 40 BIOSIS COPYRIGHT (c) 2005 The Thomson Corporation on L84 STN 1996:323333 BIOSIS ACCESSION NUMBER: PREV199699045689 DOCUMENT NUMBER: Bromoperoxidase in Corallina pilulifera is regulated by its TITLE: vanadate content. Itoh, Nobuya; Sasaki, Hiroe; Ohsawa, Noboru; Shibata, AUTHOR (S): Masaru S.; Miura, Jun'ichiro Dep. Appl. Chem. Biotechnol., Fac. Eng., Fukui Univ., CORPORATE SOURCE: Bunkyo 3-9-1, Fukui 910, Japan Phytochemistry (Oxford), (1996) Vol. 42, No. 2, pp. SOURCE: 277-281. CODEN: PYTCAS. ISSN: 0031-9422. DOCUMENT TYPE: Article LANGUAGE: English Entered STN: 11 Jul 1996 ENTRY DATE: Last Updated on STN: 11 Jul 1996 Seasonal changes in bromoperoxidase activity in coralline algae AB (Corallinaceae) are responsible for the production of volatile halogenated compounds. SDS-polyacrylamide gel electrophoresis (PAGE) of a crude protein extract showed that the concentration of this enzyme was almost constant throughout the year. Therefore, the enzyme activity in vivo changed seasonally due to a structural alteration. To elucidate this, the metal content of this enzyme at different states of activity was measured. The results revealed that the enzyme activity is controlled by the incorporation of vanadate ions, less than 1.2 mol enzyme-1, in the active site of the enzyme. Major Concepts IT Biochemistry and Molecular Biophysics; Climatology (Environmental Sciences); Enzymology (Biochemistry and Molecular Biophysics); Metabolism Chemicals & Biochemicals ITBROMOPEROXIDASE; VANADATE; VANADIUM; IRON Miscellaneous Descriptors IT ENZYME ACTIVITY; IRON; METAL ENZYME; SEASONAL CHANGE; STRUCTURAL ALTERATIONS; VANADIUM; VOLATILE HALOGENATED COMPOUND PRODUCTION ORGN Classifier

Rhodophyta 14700

Super Taxa

Algae; Plantae

Organism Name

Corallina pilulifera

Taxa Notes

Algae, Microorganisms, Nonvascular Plants, Plants

RN69279-19-2 (BROMOPEROXIDASE)

37353-31-4 (VANADATE)

7440-62-2 (VANADIUM)

7439-89-6 (IRON)

L84 ANSWER 39 OF 40 BIOSIS COPYRIGHT (c) 2005 The Thomson Corporation on

STN

ACCESSION NUMBER: 1995:159566 BIOSIS DOCUMENT NUMBER: PREV199598173866

P53 Quantitative immunocytochemical analysis in breast TITLE:

carcinomas.

AUTHOR (S): Charpin, Colette [Reprint author]; Devictor, Benedicte;

Andrac, Lucile; Amabile, Joelle; Bergeret, Denise; Lavaut,

Marie-Noelle; Allasia, Claude; Piana, Lucien

CORPORATE SOURCE: Laboratoire d'Anatomie Pathologique, Faculte de Medecine

Timone, 27 Blvd Jean Moulin, 13385 Marseielle Cedex V,

France

SOURCE: Human Pathology, (1995) Vol. 26, No. 2, pp. 159-166.

CODEN: HPCQA4. ISSN: 0046-8177.

DOCUMENT TYPE: Article English LANGUAGE:

Entered STN: 11 Apr 1995 ENTRY DATE:

Last Updated on STN: 12 Apr 1995

A series of 200 breast carcinomas was investigated on frozen sections AΒ using PAb 1801 p53 monoclonal antibody and streptavidin biotin peroxidase complex. Densitometric analysis of the immunoprecipitates was assessed by processing digitized microscopic images. p53 was observed in the nucleus of 48% of the tumors. Some tumors (14 of 91) tested in parallel on paraffin sections were negative, although positive on frozen sections. Image analysis showed that the surfaces positive with anti-p53 and the staining intensity were decreased (P lt .01) on paraffin sections. The p53 tumor expression was independent of patient age, tumor size, axillary lymph node status, HER-2/neu and cathepsin D expression, and nuclear morphometric parameters. However, p53 correlated with high histological grade (P lt .01), lack of estrogen receptor (ER) (P = .0015) and progesterone (PR) (P = .0065) antigenic sites, pS2 detection (P = .03), high Ki-67 immunoreactivity (P = .018), large silver-stained nucleolar organizer region (AgNOR) nuclear surface ratio (P lt .02), and degree of hyperploidy (P lt .03), and was more often observed in the comedocarcinomas. The results suggest that p53 expression in breast carcinomas is not a totally independent prognostic indicator and that the clinical relevance and prognostic significance of p53 expression in breast carcinomas can be reliably assessed provided that the procedures are standardized, particularly with regard to the use of frozen sections and image analysis processing of the immunodetection.

IT Major Concepts

Cell Biology; Endocrine System (Chemical Coordination and Homeostasis); Genetics; Oncology (Human Medicine, Medical Sciences); Pathology

Chemicals & Biochemicals IT

PROGESTERONE

Miscellaneous Descriptors IT

ANALYTICAL METHOD; DIAGNOSTIC METHOD; ESTROGEN; PROGESTERONE;

SINGLE-STRAND CONFORMATION POLYMORPHISM

ORGN Classifier

Hominidae 86215

Super Taxa

Primates; Mammalia; Vertebrata; Chordata; Animalia

Organism Name

human

Taxa Notes

Animals, Chordates, Humans, Mammals, Primates, Vertebrates

RN 57-83-0 (PROGESTERONE)

L84 ANSWER 40 OF 40 BIOSIS COPYRIGHT (c) 2005 The Thomson Corporation on

STN

ACCESSION NUMBER: 1994:183665 BIOSIS DOCUMENT NUMBER: PREV199497196665

TITLE: Characterization of two cold-sensitive mutants of the

beta-galactosidase from Lactobacillus delbrueckii subsp.

bulgaricus.

AUTHOR(S): Adams, Robin M.; Yoast, Sienna; Mainzer, Stanley E.; Moon,

Keith; Palombella, Anthony L.; Estell, David A.; Power,

Scott D.; Schmidt, Brian F. [Reprint author]

CORPORATE SOURCE: Khepri Pharmaceuticals Inc., 260 Littlefield Ave., South

San Francisco, CA 94080, USA

SOURCE: Journal of Biological Chemistry, (1994) Vol. 269, No. 8,

pp. 5666-5672.

CODEN: JBCHA3. ISSN: 0021-9258.

DOCUMENT TYPE: Article LANGUAGE: English

ENTRY DATE: Entered STN: 26 Apr 1994

Last Updated on STN: 18 Nov 1994

Methoxylamine mutagenesis of the beta-galactosidase gene from ΔR Lactobacillus delbruckii subsp. bulgaricus was used to generate cold-sensitive variants. Two variants, P429S and L317F, were characterized kinetically in order to determine the enzymatic consequences of these mutations. The kinetic parameters K-m and V-max on the synthetic substrate o-nitrophenyl-beta-D-galactopyranoside have been determined over a temperature range of 11-45 degree C. Only the V-max of the two variants was significantly different than the wild-type enzyme over the temperature range studied. The V-max of the L317F variant is reduced proportionately at all temperatures compared to the wild-type enzyme while the value of V-max for the P429S mutant deviates from wild-type only at lower temperatures (in 2 mM Mg-2+). This temperature-dependent effect on the V-max of P429S can be suppressed by increasing the Mg-2+ concentration. The results suggest that the binding of this essential metal ion is altered in the P429S variant such that its dissociation is increased by lowering the temperature.

IT Major Concepts

Biochemistry and Molecular Biophysics; Enzymology (Biochemistry and Molecular Biophysics); Genetics; Physiology

IT Chemicals & Biochemicals

BETA-GALACTOSIDASE; METHOXYLAMINE; O-NITROPHENYL-BETA-D-GALACTOPYRANOSIDE

IT Miscellaneous Descriptors

ENZYME KINETICS; ESSENTIAL METAL ION BINDING

ALTERATION; MAGNESIUM ION BINDING; METHOXYLAMINE; MUTAGEN; O-NITROPHENYL-BETA-D-GALACTOPYRANOSIDE; SYNTHETIC SUBSTRATE

ORGN Classifier

Regular Nonsporing Gram-Positive Rods 07830

Super Taxa

Eubacteria; Bacteria; Microorganisms

Organism Name

regular nonsporing gram-positive rods Lactobacillus delbrueckii ssp. bulgaricus

RN

Bacteria, Eubacteria, Microorganisms

9031-11-2 (BETA-GALACTOSIDASE)

67-62-9 (METHOXYLAMINE)

369-07-3 (O-NITROPHENYL-BETA-D-GALACTOPYRANOSIDE)

L85 ANSWER 1 OF 21 CAPLUS COPYRIGHT 2005 ACS on STN DUPLICATE 1

2003:216696 CAPLUS ACCESSION NUMBER:

139:113651 DOCUMENT NUMBER:

"Plugging into Enzymes": Nanowiring of Redox TITLE:

Enzymes by a Gold Nanoparticle

AUTHOR (S): Xiao, Yi; Patolsky, Fernando; Katz, Eugenii;

Hainfeld, James F.; Willner, Itamar

Institute of Chemistry, The Hebrew University of CORPORATE SOURCE:

Jerusalem, Jerusalem, 91904, Israel

Science (Washington, DC, United States) (2003), SOURCE:

299(5614), 1877-1881

CODEN: SCIEAS; ISSN: 0036-8075

American Association for the Advancement of Science PUBLISHER:

DOCUMENT TYPE: Journal English LANGUAGE:

REFERENCE COUNT: 19 THERE ARE 19 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L85 ANSWER 2 OF 21 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

2005:535294 CAPLUS

DOCUMENT NUMBER:

143:168994

TITLE: Enzymatic control of metal

deposition as key step for a low-background

electrical detection for DNA chips

Moeller, Robert; Powell, Richard D.; Hainfeld, AUTHOR (S):

James F.; Fritzsche, Wolfgang

CORPORATE SOURCE: Institute for Physical High Technology, Jena, 07702, Germany

Nano Letters (2005), 5(7), 1475-1482 SOURCE:

CODEN: NALEFD; ISSN: 1530-6984

PUBLISHER:

American Chemical Society

DOCUMENT TYPE: LANGUAGE:

Journal English

REFERENCE COUNT:

THERE ARE 32 CITED REFERENCES AVAILABLE FOR THIS 32

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L85 ANSWER 3 OF 21 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

2005:125538 CAPLUS

DOCUMENT NUMBER:

142:274639

TITLE:

Gold- and silver-facilitated metallographic in situ

hybridization procedures for detection of HER2 gene

amplification

AUTHOR (S):

Tubbs, Raymond R.; Pettay, James; Skacel, Marek; Downs-Kelly, Erinn; Powell, Richard D.; Hicks, David

G.; Hainfeld, James F.

CORPORATE SOURCE:

Department of Clinical Pathology and the Cleveland Clinic, Lerner College of Medicine, The Cleveland

Clinic Foundation, Cleveland, OH, USA

SOURCE: Molecular Morphology in Human Tissues (2005), 101-106.

Editor(s): Hacker, Gerhard W.; Tubbs, Raymond R. CRC

Press LLC: Boca Raton, Fla.

CODEN: 69GLWK; ISBN: 0-8493-1702-9

DOCUMENT TYPE: Conference; General Review

LANGUAGE: English

REFERENCE COUNT: 22 THERE ARE 22 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L85 ANSWER 4 OF 21 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2005:125537 CAPLUS

DOCUMENT NUMBER: 142:332353

TITLE: Gold cluster labels and related technologies in

molecular morphology

AUTHOR(S): Hainfeld, James F.; Powell, Richard D.

CORPORATE SOURCE: Department of Biology, Brookhaven National Laboratory,

Upton, NY, USA

SOURCE: Molecular Morphology in Human Tissues (2005), 81-100.

Editor(s): Hacker, Gerhard W.; Tubbs, Raymond R. CRC

Press LLC: Boca Raton, Fla.

CODEN: 69GLWK; ISBN: 0-8493-1702-9

DOCUMENT TYPE: Conference LANGUAGE: English

REFERENCE COUNT: 67 THERE ARE 67 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L85 ANSWER 5 OF 21 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2004:1119356 CAPLUS

DOCUMENT NUMBER: 142:476560

TITLE: Novel bright field molecular morphology methods for

detection of HER2 gene amplification

AUTHOR(S): Tubbs, Raymond; Pettay, James; Hicks, David; Skacel,

Marek; Powell, Richard; Grogan, Tom; Hainfeld,

James

CORPORATE SOURCE: Departments of Anatomic and Clinical Pathology, The

Cleveland Clinic Foundation and The Cleveland Clinic Lerner College of Medicine, Cleveland, OH, 44195, USA

SOURCE: Journal of Molecular Histology (2004), 35(6), 589-594

CODEN: JMHOAO; ISSN: 1567-2379

PUBLISHER: Kluwer Academic Publishers DOCUMENT TYPE: Journal; General Review

LANGUAGE: English

REFERENCE COUNT: 33 THERE ARE 33 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L85 ANSWER 6 OF 21 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2002:420085 CAPLUS

DOCUMENT NUMBER: 138:118072

TITLE: Supersensitive in situ hybridization by tyramide

signal amplification and nanogold silver staining: The contribution of autometallography and catalyzed reporter deposition to the rejuvenation of in situ

hybridization

AUTHOR(S): Tubbs, Raymond R.; Pettay, James; Grogan, Thomas;

Cheung, Annie L. M.; Powell, Richard D.; Hainfeld, James; Hauser-Kronberger, Cornelia;

Hacker, Gerhard W.

CORPORATE SOURCE: Department of Clinical Pathology, The Cleveland Clinic

Foundation, Cleveland, OH, USA

SOURCE: Gold and Silver Staining (2002), 127-144. Editor(s):

Page 56 12/05/2005 Searched by Alex Waclawiw

Hacker, Gerhard W.; Gu, Jiang. CRC Press LLC: Boca

Raton, Fla.

CODEN: 69CQTC; ISBN: 0-8493-1392-9

DOCUMENT TYPE: LANGUAGE:

Conference English

REFERENCE COUNT:

25 THERE ARE 25 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L85 ANSWER 7 OF 21 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

2000:658520 CAPLUS

DOCUMENT NUMBER:

133:251273

TITLE:

Metal-lipid molecules

INVENTOR(S):

Hainfeld, James F.; Furuya, Frederic R.;

Powell, Richard D.; Joshi, Vishwas N.; Gutierrez,

Edmund

PATENT ASSIGNEE(S):

Nanoprobes, Inc., USA

SOURCE:

U.S., 20 pp., Cont.-in-part of U.S. 5,728,590.

CODEN: USXXAM

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 6121425	Α	20000919	US 1998-39601	19980316
US 5521289	A	19960528	US 1994-282929	19940729
US 5728590	A	19980317	US 1996-652007	19960523
US 6369206	B1	20020409	US 2000-619343	20000719
US 6818199	B1	20041116	US 2002-93770	20020308
US 2005130207	A1	20050616	US 2004-988781	20041115
PRIORITY APPLN. INFO.:			US 1994-282929	A2 19940729
			US 1996-652007	A2 19960523
			US 1998-39601	A1 19980316
			US 2000-619343	A2 20000719
			US 2002-93770	A1 20020308

REFERENCE COUNT: 19 THERE ARE 19 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L85 ANSWER 8 OF 21 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

1999:668511 CAPLUS

DOCUMENT NUMBER:

132:262328 Gold-ATP

TITLE: AUTHOR(S):

Tologold Tomos D. Tile

Montgerra

CORPORATE SOURCE:

Hainfeld, James F.; Liu, Wenqiu; Barcena,
Montserrat

Biology Department, Brookhaven National Laboratory,

SOURCE:

Upton, NY, 11973, USA Journal of Structural Biology (1999), 127(2), 120-134

CODEN: JSBIEM; ISSN: 1047-8477

PUBLISHER:

Academic Press

DOCUMENT TYPE:

Journal English

LANGUAGE:
REFERENCE COUNT:

61 THERE ARE 61 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L85 ANSWER 9 OF 21 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

1985:200240 CAPLUS

DOCUMENT NUMBER:

102:200240

TITLE:

Escherichia coli pyruvate dehydrogenase complex: particle masses of the complex and component

Page 57 12/05/2005 Searched by Alex Waclawiw

enzymes measured by scanning transmission

electron microscopy

CaJacob, Claire A.; Frey, Perry A.; Hainfeld, AUTHOR (S):

James F.; Wall, Joseph S.; Yang, Heechung

CORPORATE SOURCE: Coll. Agric. Life Sci., Univ. Wisconsin, Madison, WI,

53705, USA

Biochemistry (1985), 24(10), 2425-31 SOURCE:

CODEN: BICHAW; ISSN: 0006-2960

DOCUMENT TYPE: Journal English LANGUAGE:

L85 ANSWER 10 OF 21 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1985:181471 CAPLUS

DOCUMENT NUMBER: 102:181471

Evidence for two spatially distinct domains on each TITLE:

subunit of methylenetetrahydrofolate reductase Matthews, Rowena G.; Vanoni, Maria A.; Khani,

Shahrokh; Hainfeld, James F.; Wall, Joseph CORPORATE SOURCE: Dep. Biol. Chem., Univ. Michigan, Ann Arbor, MI,

48109, USA

Flavins Flavoproteins, Proc. Int. Symp., 8th (1984), SOURCE:

> 217-20. Editor(s): Bray, Robert C.; Engel, Paul C.; Mayhew, Stephen G. de Gruyter: Berlin, Fed. Rep. Ger.

CODEN: 53HMAL

Conference DOCUMENT TYPE: LANGUAGE: English

L85 ANSWER 11 OF 21 BIOSIS COPYRIGHT (c) 2005 The Thomson Corporation on

STN

AUTHOR (S):

ACCESSION NUMBER: 2005:521812 BIOSIS DOCUMENT NUMBER: PREV200510292858

Analytical validation and interobserver reproducibility of TITLE:

EnzMet GenePro - A second-generation bright-field

metallography assay for concomitant detection of HER2 gene status and protein expression in invasive carcinoma of the

breast.

Downs-Kelly, Erinn; Pettay, James; Hicks, David; Skacel, AUTHOR (S):

Marek; Yoder, Brian; Rybicki, Lisa; Myles, Jonathan; Sreenan, Joseph; Roche, Patrick; Powell, Richard; Hainfeld, James; Grogan, Thomas; Tubbs, Raymond

[Reprint Author]

CORPORATE SOURCE: 9500 Euclid Ave, L11, Cleveland, OH 44195 USA

TubbsR@ccf.org

American Journal of Surgical Pathology, (NOV 2005) Vol. 29, SOURCE:

No. 11, pp. 1505-1511.

ISSN: 0147-5185.

DOCUMENT TYPE:

Article English

LANGUAGE:

Entered STN: 23 Nov 2005 ENTRY DATE:

Last Updated on STN: 23 Nov 2005

L85 ANSWER 12 OF 21 BIOSIS COPYRIGHT (c) 2005 The Thomson Corporation on

STN

ACCESSION NUMBER: 2005:415036 BIOSIS DOCUMENT NUMBER: PREV200510210247

Enzyme metallography (EnzMet) - A robust TITLE:

detection system for high resolution ultrasensitive

immunohistochemistry (IHC).

Tubbs, R. R. [Reprint Author]; Pettay, J.; Roche, P. C.; AUTHOR (S):

Powell, W.; Powell, R. D.; Grogan, T.; Hainfeld, J.

F

CORPORATE SOURCE:

Cleveland Clin Fdn, Cleveland, OH 44195 USA

SOURCE:

Modern Pathology, (JAN 2005) Vol. 18, No. Suppl. 1, pp.

335A.

Meeting Info.: 94th Annual Meeting of the

United-States-and-Canadian-Academy-of-Pathology. San

Antonio, TX, USA. February 26 -March 04, 2005. US Canadian

Acad Pathol. ISSN: 0893-3952.

DOCUMENT TYPE:

Conference; (Meeting)

Conference; Abstract; (Meeting Abstract)

LANGUAGE:

English

ENTRY DATE:

Entered STN: 19 Oct 2005

Last Updated on STN: 19 Oct 2005

L85 ANSWER 13 OF 21 BIOSIS COPYRIGHT (c) 2005 The Thomson Corporation on

STN

ACCESSION NUMBER:

2002:426317 BIOSIS PREV200200426317

DOCUMENT NUMBER: TITLE:

Interobserver interpretative reproducibility of GOLDFISH, a first generation gold-facilitated autometallographic bright

field in situ hybridization assay for HER-2/neu amplification in invasive mammary

carcinoma.

AUTHOR (S):

Tubbs, Raymond [Reprint author]; Skacel, Marek; Pettay, James; Powell, Richard; Myles, Jonathan; Hicks, David; Sreenan, Joseph; Roche, Patrick; Stoler, Mark H.;

Hainfeld, James

CORPORATE SOURCE:

Department of Clinical Pathology, 9500 Euclid Avenue, L-11,

Cleveland, OH, 44195-5131, USA

tubbsr@ccf.org

SOURCE:

American Journal of Surgical Pathology, (July, 2002) Vol.

26, No. 7, pp. 908-913. print.

ISSN: 0147-5185.

DOCUMENT TYPE:

LANGUAGE:

Article English

ENTRY DATE:

Entered STN: 7 Aug 2002

Last Updated on STN: 7 Aug 2002

L85 ANSWER 14 OF 21 BIOSIS COPYRIGHT (c) 2005 The Thomson Corporation on

STN

ACCESSION NUMBER:

2002:318388 BIOSIS PREV200200318388

DOCUMENT NUMBER: TITLE:

Interobserver reproducibility of a first generation

autometallographic bright field assay for Her-

2/new amplification (GOLDFISH).

AUTHOR(S):

Tubbs, R. R. [Reprint author]; Skacel, M. [Reprint author]; Pettay, J. D. [Reprint author]; Powell, R. D.; Myles, J. L. [Reprint author]; Sreenan, J. J.; Hicks, D. G. [Reprint author]; Stoler, M. H.; Roche, P. C.; Jenkins, R. B.;

Hainfeld, J. F.

CORPORATE SOURCE:

Cleveland Clinic, Cleveland, OH, USA

SOURCE:

Laboratory Investigation, (January, 2002) Vol. 82, No. 1,

pp. 54A. print.

Meeting Info.: 2002 Annual Meeting of the United States and Canadian Academy of Pathology. Chicago, IL, USA. February

23-March 01, 2002.

CODEN: LAINAW. ISSN: 0023-6837.

DOCUMENT TYPE:

Conference; (Meeting)

Conference; Abstract; (Meeting Abstract)

LANGUAGE: English

ENTRY DATE: Entered STN: 5 Jun 2002

Last Updated on STN: 5 Jun 2002

L85 ANSWER 15 OF 21 BIOSIS COPYRIGHT (c) 2005 The Thomson Corporation on

STN

ACCESSION NUMBER: 2002:318354 BIOSIS DOCUMENT NUMBER: PREV200200318354

TITLE: Validation of GOLDFISH (Gold Facilitated Autometallographic

In Situ Hybridization) for assessment of Her-

2/neu oncogene amplification by

conventional microscopy: A detailed investigation of

discrepancies with direct fluorescence in situ

hybridization (FISH).

AUTHOR(S): Pettay, J. D. [Reprint author]; Skacel, M. [Reprint

author]; Powell, R. D.; Stoler, M. H.; Roche, P. C.;

Hainfeld, J. F.; Tubbs, R. R. [Reprint author]

CORPORATE SOURCE: Cleveland Clinic, Cleveland, OH, USA

SOURCE: Laboratory Investigation, (January, 2002) Vol. 82, No. 1,

pp. 46A. print.

Meeting Info.: 2002 Annual Meeting of the United States and Canadian Academy of Pathology. Chicago, IL, USA. February

23-March 01, 2002.

CODEN: LAINAW. ISSN: 0023-6837.

DOCUMENT TYPE: Conference; (Meeting)

Conference; Abstract; (Meeting Abstract)

LANGUAGE: English

ENTRY DATE: Entered STN: 5 Jun 2002

Last Updated on STN: 5 Jun 2002

L85 ANSWER 16 OF 21 BIOSIS COPYRIGHT (c) 2005 The Thomson Corporation on

STN

ACCESSION NUMBER: 2000:221042 BIOSIS DOCUMENT NUMBER: PREV200000221042

TITLE: Introduction of a novel HRP substrate-Nanogold probe for

signal amplification in immunocytochemistry.

AUTHOR(S): Mayer, Gaetan; Leone, Robert D.; Hainfeld, James F.

; Bendayan, Moise [Reprint author]

CORPORATE SOURCE: Dept. of Pathology and Cell Biology, Universite de

Montreal, Succ. Centre Ville, Montreal, Quebec, H3C 317,

Canada

SOURCE: Journal of Histochemistry and Cytochemistry, (April, 2000)

Vol. 48, No. 4, pp. 461-469. print.

CODEN: JHCYAS. ISSN: 0022-1554.

DOCUMENT TYPE: Article LANGUAGE: English

ENTRY DATE: Entered STN: 31 May 2000

Last Updated on STN: 5 Jan 2002

L85 ANSWER 17 OF 21 BIOSIS COPYRIGHT (c) 2005 The Thomson Corporation on

STN

ACCESSION NUMBER: 1997:251852 BIOSIS DOCUMENT NUMBER: PREV199799551055

TITLE: Sensitive in situ hybridization with catalyzed reporter

deposition, streptavidin-Nanogold, and silver

acetate autometallography: Detection of single-copy human

papillomavirus.

AUTHOR(S): Zehbe, Ingeborg [Reprint author]; Hacker, Gerhard W.; Su,

Huici; Hauser-Kronberger, Cornelia; Hainfeld, James

F.; Tubbs, Raymond

Deutsches Krebsforschungszentrum, Angewandt Tumorvirol., CORPORATE SOURCE:

Virus-Wirtszell-Wechselwirkungen, Im Neuenheimer Feld 242,

D-69120 Heidelberg, Germany

American Journal of Pathology, (1997) Vol. 150, No. 5, pp. SOURCE:

1553-1561.

CODEN: AJPAA4. ISSN: 0002-9440.

DOCUMENT TYPE:

Article

LANGUAGE:

English

ENTRY DATE:

Entered STN: 13 Jun 1997

Last Updated on STN: 13 Jun 1997

L85 ANSWER 18 OF 21 BIOSIS COPYRIGHT (c) 2005 The Thomson Corporation on

ACCESSION NUMBER:

1994:448312 BIOSIS PREV199497461312

DOCUMENT NUMBER: TITLE:

Mapping the lipoyl groups of the pyruvate dehydrogenase

complex by use of gold cluster labels and

scanning transmission electron microscopy.

AUTHOR (S):

Yang, Yuh-Shyong; Datta, Asit; Hainfeld, James F.

; Furuya, Frederic R.; Wall, Joseph S.; Frey, Perry A.

[Reprint author]

CORPORATE SOURCE:

Inst. for Enzyme Res., Graduate Sch., Dep. Biochemistry,

College of Agricultural and Life Sciences, Univ.

Wisconsin-Madison, Madison, WIS 53705, USA

SOURCE:

Biochemistry, (1994) Vol. 33, No. 32, pp. 9428-9437. CODEN: BICHAW. ISSN: 0006-2960.

DOCUMENT TYPE:

Article English

LANGUAGE: ENTRY DATE:

Entered STN: 24 Oct 1994

Last Updated on STN: 25 Oct 1994

L85 ANSWER 19 OF 21 BIOSIS COPYRIGHT (c) 2005 The Thomson Corporation on

ACCESSION NUMBER:

1993:321665 BIOSIS

DOCUMENT NUMBER:

PREV199396030015

TITLE:

A polypeptide bound by the chaperonin groEL is localized

within a central cavity.

AUTHOR(S):

Braig, Kerstin [Reprint author]; Simon, Martha; Furuya,

Fred; Hainfeld, James F.; Horwich, Arthur L.

[Reprint author]

CORPORATE SOURCE:

Howard Hughes Med. Inst., Dep. Genetics, Yale Univ. Sch.

Med., 333 Cedar St., New Haven, CT 06510, USA

SOURCE:

Proceedings of the National Academy of Sciences of the United States of America, (1993) Vol. 90, No. 9, pp.

3978-3982.

CODEN: PNASA6. ISSN: 0027-8424.

DOCUMENT TYPE:

Article

LANGUAGE:

English

ENTRY DATE:

Entered STN: 12 Jul 1993

Last Updated on STN: 13 Jul 1993

L85 ANSWER 20 OF 21 BIOSIS COPYRIGHT (c) 2005 The Thomson Corporation on

STN

ACCESSION NUMBER:

1992:225044 BIOSIS

DOCUMENT NUMBER:

PREV199242106544; BR42:106544

TITLE:

ELECTRON MICROSCOPY OF UNSTAINED GOLD-CLUSTER

LABELED PYRUVATE AND 2 OXOGLUTARATE DEHYDROGENASE COMPLEXES

PDC AND OGDC.

AUTHOR (S):

WAGENKNECHT T [Reprint author]; GRASSUCCI R; BERKOWITZ J;

CARBONE K; FURUYA F; HAINFELD J

CORPORATE SOURCE: WADSWORTH CENT LAB RES, NY STATE DEP HEALTH, ALBANY, NY

12201-0509, USA

SOURCE: Biophysical Journal, (1992) Vol. 61, No. 2 PART 2, pp.

A469.

Meeting Info.: JOINT ANNUAL MEETING OF THE BIOPHYSICAL SOCIETY AND THE AMERICAN SOCIETY FOR BIOCHEMISTRY AND MOLECULAR BIOLOGY, HOUSTON, TEXAS, USA, FEBRUARY 9-13,

1992. BIOPHYS J.

CODEN: BIOJAU. ISSN: 0006-3495.

DOCUMENT TYPE:

Conference; (Meeting)

FILE SEGMENT:

BR

LANGUAGE:

ENGLISH

ENTRY DATE:

Entered STN: 5 May 1992

Last Updated on STN: 6 May 1992

L85 ANSWER 21 OF 21 BIOSIS COPYRIGHT (c) 2005 The Thomson Corporation on

STN

ACCESSION NUMBER: 1992:177621 BIOSIS

DOCUMENT NUMBER: PREV199242082621; BR42:82621

TITLE: ELECTRON MICROSCOPY OF UNSTAINED GOLD-CLUSTER

LABELED PYRUVATE AND 2 OXOGLUTARATE DEHYDROGENASE COMPLEXES

PDC AND OGDC.

AUTHOR(S): WAGENKNECHT T [Reprint author]; GRASSUCCI R; BERKOWITZ J;

CARBONE K; FURUYA F; HAINFELD J

CORPORATE SOURCE: WADSWORTH CENT LAB RES, NEW YORK STATE DEP HEALTH, ALBANY,

NY 12201, USA

SOURCE: FASEB Journal, (1992) Vol. 6, No. 1, pp. A469.

Meeting Info.: JOINT MEETING OF THE AMERICAN SOCIETY FOR BIOCHEMISTRY AND MOLECULAR BIOLOGY/BIOPHYSICAL SOCIETY, HOUSTON, TEXAS, USA, FEBRUARY 9-13, 1992. FASEB (FED AM SOC

EXP BIOL) J.

CODEN: FAJOEC. ISSN: 0892-6638.

DOCUMENT TYPE:

Conference; (Meeting)

FILE SEGMENT: H

LANGUAGE:

ENTRY DATE: Entered STN: 2 Apr 1992

ENGLISH

Last Updated on STN: 3 Apr 1992

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